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(US). WATSON, Brian, Mory Place, Carmel, IN 46033 (US).

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Inventors; and

EE

(81) Designated States (national): AB, (71) Applicant (for all designated States except US): ELI LILLY AND COMPANY [US/US]; Patent Division, P. O. Box 6288, Indianapolis, IN 46206-6288 (US). English S

3 [19] Inventoral Applicants (for US oaly): BEAVERS, Lida, (if Sebam (USUS); 191 West State Road 222, Frankin, IN 46131 (Lid); GADSKI, Robert, Alan (USUS); 443 North Illinois, Indianapolis, IN 46208 (US). HIPSKIND, Philip, Arthur (USUS); 1825 South Cabic Cont. New Palettine, IN 46143 (US). LIMDSIEX, Crafg, William (USUS); 126 Berger Road, Schwenszwille, PA 19473

(54) TIHE: NON-IMIDAZOLE ARYL, ALKYLAMINES COMPOUNDS AS HISTAMINE H3 RECEPTOR ANTAGONISTS. PREPARATION AND THERAPEUTIC USES

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Abstract. The present invention discloses novel substituted any altylamine compounds of Formula (0) or pharmaceutically quable satis thereofwhich have selective histornic-H3 recaptors anagonist activity as well as methods for preparing such commiss. In another embodiment, the invention discloses pharmaceutical compositions comprising such cyclic amines as well as: pounds. In another embodiment, the invention discloses pharmaceutical compositions methods of using them to treat obesity and other histamine H3 receptor-related disease (57) Abstract: The present 7Y S769L0/70 OM

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NON-IMIDAZOLE ARYL ALKYLAMINES COMPOUNDS AS HISTAMINE H3 RECEPTOR ANTAGONISTS, PREPARATION AND THERAPEUTIC USES

BACKGROUND OF THE INVENTION

The present invention relates to histamine H3 receptor antagonists, and as such are receptors, such as obesity, cognitive disorders, attention deficient disorders and the like. useful in the treatment of disorders responsive to the inactivation of histamine H3

nistamine H3 receptor is relatively neuron specific and inhibits the release of a number of receptor increase synthesis and release of cerebral histamine and other monoamines. By mood and attention adjustments, cognitive deficiencies, obesity, dizziness, schizophrenis receptor found in the peripheral and central nervous system and regulates the release of histamine H3 receptor is an important target for new therapeutics in Alzheimer disease, monamines, including histamine. Selective antagonism of the histamine H3 receptor minimizing non-specific peripheral consequences. Antagonists of the histamine H3 this mechanism, they induce a prolonged wakefulness, improved cognitive function, reduction in food intake and normalization of vestibular reflexes. Accordingly, the raises brain histamine levels and inhibits such activities as food consumption while The histamine H3 receptor (H3R) is a presynaptic autoreceptor and heteronistamine and other neurotransmitters, such as serotonin and acetylcholine. The epilepsy, sleeping disorders, narcolepsy and motion sickness. 2 2

The majority of histamine H3 receptor antagonists to date resemble histamine in 494010, WO 97/29092, WO 96/38141, and WO96/38142. These imidazole-containing possessing an imidazole ring generally substituted in the 4(5) position (Ganellin et al., Ars Pharmaceutica, 1995, 36:3, 455-468). A variety of patents and patent applications compounds have the disadvantage of poor blood-brain barrier penetration, interaction directed to antagonists and agonists having such structures include EP 197840, EP with cytochrome P-450 proteins, and hepatic and ocular toxicities.

23

Pharm. 1985, 111:72-84) demonstrated some histamine H3 receptor activity but with poor Non-imidazole neuroactive compounds such as beta histamines (Arrang, Eur. J. potency. EP 978512 published March 1, 2000 discloses non-imidazole aryloxy

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alkylamines discloses histamine H3 receptor antagonists but does not disclose the affinity, of a saturated, fused heterocyclic ring appended to the central benzene core. Furthermore if any, of these antagonists for recently identified histamine receptor GPRv53, described substitutions of the non-oxygen benzene ring substituent, and in some cases the presence substitutions at the ortho, meta or para positions of the central benzene ring, the exact below. EP 0982300A2 (pub. March 1, 2000) discloses non-imidazole alkyamines as histamine HS receptor ligand which are similar to the subject invention by having a the compounds of this invention are highly selective for the H3 receptor (vs. other phenoxy core structure although the subject invention is unique in the dissimilar

Histamine mediates its activity via four receptor subtypes, H1R, H2R, H3R and a histamine receptors), and possess remarkable drug disposition properties (pharmacokinetics) 2

newly identified receptor designated GPRv53 [(Oda T., et al., J.Biol.Chem. 275 (47):

effects when targeting antagonism of the H3R receptor. Furthermore, the identification of H2R and H3R, few specific ligands have been developed that can distinguish H3R from this new receptor has fundamentally changed histamine biology and must be considered 36781-6 (2000)]. Although relatively selective ligands have been developed for H1R, leukocytes. Activation or inhibition of this receptor could result in undesirable side GPRv53. GPRv53 is a widely distributed receptor found at high levels in human 2

Because of the unresolved deficiencies of the compounds described above, there is a continuing need for improved methods and compositions to treat disorders associated with histamine H3 receptors.

in the development of histamine H3 receptor antagonists.

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receptor antagonists. In another aspect, the present invention provides compounds that are useful as selective antagonists of the histamine H3 receptor but have little or no The present invention provides compounds that are useful as histamine H3 pharmaceutical compositions comprising antagonists of the histamine H3 receptor binding affinity of GPRv53. In yet another aspect, the present invention provides

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In yet another aspect, the present invention provides compounds, pharmaceutical attention deficient disorders and other disorders associated with histamine H3 receptor compositions, and methods useful in the treatment of obesity, cognitive disorders, 30

SUMMARY OF THE INVENTION

The present invention is a compound structurally represented by Formula I

or pharmaceutically acceptable salts thereof wherein:

X is O, NR7 or S;

10 R¹ is hydrogen,

 $C_1\hbox{-} C_8$ alkyl optionally substituted with 1 to 4 halogens,

(CHR5)n-C3-C7 cycloalkyl,

(CHR⁵)_h aryl,

(CHR⁵)_h heteroaryl, or

 $(CHR^5)_n$ -O $(CHR^5)_n$ -aryl;

15

R2 is independently R1, or

 COR^1 , or cyclized with the attached nitrogen atom at the R^1 position to form a 4, 5, or 6 member carbon ring, wherein one of said carbons is optionally replaced by one of

O, S, NR 1 or CO, or wherein the ring formed by R 1 and R 2 is optionally substituted one to two times with C1-C4 alkyl;

R³ is independently C₃-C₇ cycloalkylene, or C₁- C₄ alkylene optionally substituted;

R4 is hydrogen,

C₁-C₄ alkyl,

(CHR⁵)_n-C₃-C₇ cycloalkyl,

(CHR⁵)_n aryl,

(CHR5)_n heteroaryl,

(CHR5)_n-O(CHR5)_n-aryl or

CO or

cyclized with R5 to from a cyclopropyl ring;

2

R⁵ is hydrogen, or

C₁-C₄ alkyl;

R⁶ is hydrogen,

halo or

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cyclized with the attached carbon atom at the $\ensuremath{R^{5}}$ position to form a 5 to 6 member

carbon ring,

cyclized with the attached carbon atom at the \mathbb{R}^7 position to form a 5 to 6 member heterocyclic ring or

2

 \mathbb{R}^7 is hydrogen,

C1-C8 alkyl optionally substituted with 1 to 4 halogens,

(CHR5)n-C3-C7 cycloalkyi,

(CHR⁵)_n aryl,

 $(CHR^5)_n$ heteroaryl,

52

(CHR⁵)_n-O(CHR⁵)_n-aryl,

SO2R1 or

Cyclized with attached carbon on R ⁸ to from a 5, 6, or 7 membered carbon ring	•
onally substituted with R9, CF3, or CN, optionally one of the said carbons is replaced	
I, NR ¹ , CO;	

by N.

C₁-C₈ alkyl -SO₂ R⁹. -CO₂ R¹⁰, 5 R⁸ is hydrogen, a bond,

-CONH R 10; -co R9, 2

R⁹ is hydrogen,

 $\mathsf{C}_1\text{-}\mathsf{C}_8$ alkyl optionally substituted with 1 to 4 halogens, C3-C7 cycloalkyl, halogen, 15

heterocycle, heteroaryl,

CH₂ aryl,

-0(CHR⁵)_n-aryl,

-CONR 1 R2,

-SO2R1,

-coR1

-OR1,

-NR¹ R², -CH₂NR¹ R², $-N(R^1)_2$,

-CONR¹ R² $-SO_2N(\mathbb{R}^1)_2$, -NHSO2R1, -CO2R1, -NO₂,

 $-S(O)_nR^1$, -CH2SR3, -0CF3,

R¹⁰ is hydrogen, halogen, 2

C₁-C₈ alkyl optionally substituted with 1 to 4 halogens, C₃-C₇ cycloalkyl,

CH₂ aryl,

heterocycle, heteroaryl,

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-cor¹, -conr¹ r²,

-SO2R1,

 $-N(R^1)_2$,

-CH2NR1 R2, -NR1 R2,

-CONR1 R2

-co₂R¹,

 $-50_2N(R^1)_2$,

-CH2SR3,

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and n is 0 - 4.

In preferred embodiments of Formula I the core phenoxy ring is an o, m, or p-disubstituted benzene, more preferably a p-disubstituted benzene. In alternative embodiments R⁶ forms a bicyclic carbon ring at the R⁵ position. Alternatively, R⁶ may form a bicyclic heterocyclic ring at the R⁷ position. Preferably, X is nitrogen, R⁴ and R⁵ are independently H or CH₅, R1 and R2 are independently a C₁-C₈ alkyl and R9 is a di-C₁ to C₂ alkyl-amino.

The present invention is a pharmaceutical composition which comprises a compound of Formula I and a pharmaceutically acceptable carrier. Pharmaceutical formulations of Formula I can provide a method of selectively increasing histamine levels in cells by contacting the cells with an antagonist of the histamine H3 receptor, the antagonists being a compound of Formula I.

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The present invention further provides an antagonist of Formula I which is characterized by having little or no binding affinity for the histamine receptor GPRv53. Thus, a pharmaceutical preparation of Formula I can be useful in the treatment or prevention of obesity, cognitive disorders, attention deficient disorders and the like, which comprises administering to a subject in need of such treatment or prevention an effective amount of a compound of Formula I. In addition, a pharmaceutical preparation of Formula I can be useful in the treatment or prevention of a disorder or disease in which inhibition of the histamine H3 receptor has a beneficial effect or the treatment or prevention of eating disorders which comprises administering to a subject in need of such treatment or prevention an effective amount of a compound of Formula I.

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DETAILED DESCRIPTION OF THE INVENTION

Throughout the instant application, the following terms have the indicated meanings:

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The term "GPRv53" means a recently identified novel histamine receptor as described in Oda, et al., supra. Alternative names for this receptor are PORT3 or H4R.

The term "H3R" means to the histamine H3 receptor that inhibits the release of a

30 number of monoamines, including histamine.

The term "H1R" means to the histamine H1 receptor subtype. The term "H2R" means to the histamine H2 receptor subtype.

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The term "selective H3R antagonists" is defined as the ability of a compound of the present invention to block forskolin-stimulated cAMP production in response to agonist $R(\cdot)\alpha$ methylhistamine.

agoniss in 1750 metry missantime.
"Alkylene" are a saturated hydrocarbyldiyl radical of straight or branched

configuration made up of from 1 to 4 carbon atoms. Included within the scope of this term are methylene, 1,2 -ethane-diyl, 1,1-ethane-diyl, 1,3-propane diyl, 1,2-propane diyl, 1,3 butane-diyl, 1,4 -butane diyl, and the like.

"C₁-C₇ cycloalkylene" are a saturated hydrocarbyldiyl radical of cyclic configuration, optionally branched, made up of from 3 to 7 carbon atoms. Included within the scope of this term are cyclopropyl, cyclobutyl, cyclopentyl and cyclohexyl, and

2

"Alkyl" are one to four or one to eight carbon atoms such as methyl, ethyl, propyl, butyl, pentyl, hexyl, heptyl, octyl and isomenc forms thereof.

"Aryl" are six to twelve carbon atoms such as phenyl, alpha -naphthyl, beta - 15 naphthyl, m-methylphenyl, p-trifluoromethylphenyl and the like. The aryl groups can also be substituted with one to 3 hydroxy, fluoro, chloro, or bromo groups.

"Cycloalkyl" are three to seven carbon atoms such as cyclopropyl, cyclobutyl, cyclopentyl and cyclohexyl.

- "Heteroaryl" are six to twelve carbon atoms aryls, as described above, containing the heteroatoms nitrogen, sulfur or oxygen. Heteroaryls are pyridine, thiophene, furan, pyrimidine, 2-pyridyl, 3-pyridyl, 2-pyrimidinyl, 4-pyrimidinyl, 5-pyrimidinyl, 3-pyrimidinyl, 2-pyrimidinyl, 1-isoquinolyl, 1-isoquinolyl, 3-pyrimidinyl, 1-isoquinolyl, 2-quinazolinyl, 4-quinazolinyl, 4-quinazolinyl, 2-quinoxalinyl, 1-phthalazinyl, 2-imidazolyl, 4-imidazolyl, 3-isoxazolyl, 4-isoxazolyl, 3-pyrazolyl, 4-
 - 25 pyrazolyl, 5-pyrazolyl, 2-oxazolyl, 4-oxazolyl, 5-oxazolyl, 2-thiazolyl, 4-thiazolyl, 5-thiazolyl, 2-indolyl, 3-indolyl, 3-indazolyl, 2-benzoxazolyl, 2-benzotimiazolyl, 2-benzofuranyl, 3-benzofuranyl, 3-furanyl, 3-turanyl, 3-turanyl, 3-turanyl, 3-turanyl, 3-turanyl, 1.2,4-oxadiazol-5-yl, 1.2,4-triazol-5-yl, 1.2,4-triazol-5-yl, 1.2,4-triazol-5-yl, 1.2,3-thiadiazol-5-yl, 1.2,4-triazol-5-yl, 1.2,3-triazol-5-yl, 1.2,3-triazol-5-yl, 1.2,4-triazol-5-yl, 1.2,3-triazol-5-yl, 1.2,3-tr
- 30 tetrazol-5-yl, 5-oxazolyl, 1-pyrrolyl, 1-pyrazolyl, 1,2,3-triazol-1-yl, 1,2,4-triazol-1-yl, 1-tetrazolyl, 1-indolyl, 1-indazolyl, 2-isoindolyl, 1-purinyl, 3-isothiazolyl, 4-isothiazolyl, 5-isonhiazolyl

'Heterocycle" are three to twelve carbon atom cyclic aliphatic rings, wherein one or more carbon atoms is replaced by a hetero-atom which is nitrogen, sulfur or oxygen.

"Halogen" or "halo" means fluoro, chloro, bromo and iodo.

Composition" means a pharmaceutical composition and is intended to encompass ingredient(s) that make up the carrier. Accordingly, the pharmaceutical compositions of the present invention encompass any composition made by admixing a compound of the a pharmaceutical product comprising the active ingredient(s), Formula I, and the inert present invention and a pharmaceutically acceptable carrier.

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The term "unit dosage form" means physically discrete units suitable as unitary predetermined quantity of active material calculated to produce the desired therapeutic dosages for human subjects and other non-human animals, each unit containing a effect, in association with a suitable pharmaceutical carrier. 2

The terms "treating" and "treat", as used herein, include their generally accepted stopping, or reversing the progression or severity of a pathological condition, described meanings, i.e., preventing, prohibiting, restraining, alleviating, ameliorating, slowing, herein.

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wherein R6 is cyclized with the attached carbon atom at R7 to form, including the fused compound wherein X is nitrogen, and wherein \mathbb{R}^7 and \mathbb{R}^8 are cyclized to form, together In one embodiment, the present invention provides compounds of Formula I benzene ring, a substituted tetrahydroisoquinoline ring. Another embodiment is a structure is an o, m, or p- disubstituted aryl. Another embodiment is a compound as described in detail above. Another embodiments are where the phenoxy core with X, a pyrrolidine ring, and wherein \mathbb{R}^9 is -CH2-N-pyrrolidinyl.

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A preferred moiety for X is independently O or N.

A preferred moiety for R9 is C1-C8 dialkylamino. A more preferred embodiment is where the dialkylamino is dimethylamino. 52

I are meant to also include the pharmaceutical salts, its enantiomers and racernic mixtures It will be understood that, as used herein, references to the compounds of Formula

Because certain compounds of the invention contain a basic moiety (e.g., amino), the compound of Formula I can exist as a pharmaceutical acid addition salt. Such salts include sulfate, pyresulfate, bisulfate, sulfite, bisulfite, phosphate, mono-

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bromide, iodide, acetate, propionate, decanoate, caprylate, acrylate, formate, isobutyrate, hydrogenphosphate, dihydrogenphosphate, metaphosphate, pyrophosphate, chloride, heptanoate, propiolate, oxalate, malonate, succinate, suberate, sebacate, fumarate, maleate, 2-butyne-1,4 dioate, 3-hexyne-2, 5-dioate, benzoate, chlorobenzoate,

glycollate, maleate, tartrate, methanesulfonate, propanesulfonate, naphthalene-i-sulfonate, phenylpropionate, phenylbutyrate, citrate, lactate, hippurate, beta-hydroxybutyrate, hydroxybenzoate, methoxybenzoate, phthalate, xylenesulfonate, phenylacetate, naphthalene-2-sulfonate, mandelate and the like salts.

stereoisomers of the compounds also. Thus, as one skilled in the art knows, certain aryls may exist in tautomeric forms. Such variations are contemplated to be within the scope As stated earlier, the invention includes tautomers, enantiomers and other of the invention. 2

The compounds of Formula I may be prepared by several processes well known in Mitsunobu chemistries and reductive animations known to one skilled in the art, or by the temperatures in the range about 0-1000 C., by bringing together the ingredients in contact hydrocarbons, toluene, acetonitrile and the like, preferably in the absence of moisture, at the art. The compounds of the present invention are prepared by standard alkylation or methods provided herein, supplemented by methods known in the art. Generally, this in the solvent medium and stirring for about 10 minutes to about 48 hours at such reaction is conducted in an organic solvent such as, for example, halogenated

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crystallization from a suitable solvent, for example methanol or ethyl acetate or a mixture The compounds of Formula I, when existing as a diastereomeric mixture, may be separated into diastereomenic pairs of enantiomers by, for example, fractional

- as a resolving agent. Alternatively, any enantiomer of a compound of the formula may be obtained by stereospecific synthesis using optically pure starting materials or reagents of stereoisomers by conventional means, for example by the use of an optically active acid thereof. The pair of enantiomers thus obtained may be separated into individual known configuration or through enantioselective synthesis. 52
- The Examples shown in Table 1 below are being provided to further illustrate the present invention. They are for illustrative purposes only; the scope of the invention is 8

not to be considered limited in any way thereby. The preparation of compounds of Formula I, are depicted in the schemes and procedures below.

Preparation of N-(1-[4-(3-Dimethylamino-propoxy)-phenyl-N', N'-dimethyl-ethane-1, 2diamine

To a 100 mL round-bottom flask was placed NaH (60% dispersion, 38.4 mg, 1.0 Example 2

DMF solution of p-hydroxyacetophenone (62 mg, 0.5 mmol) was added at 0 C. After 15 The reaction was then quenched with water, diluted with ether and washed with water (3 was added, and the reaction was allowed to slowly reach room temperature over 3 hours. x 20 mL) and brine (2x 20 mL). Concentration in vacuo afforded 114 mg (92%) of an minutes, a DMF solution of 3-chloro-N,N-diethyl-N-proplyamine (150 mg, 1.0 mmol) mmol) and anhydrous THF (10 mL, 0.1 M) under an atmosphere of nitrogen. Then, a 2

off-white solid. LCMS indicated a purity of 95% and hit the mass, 249.1. This material temperature, NaCNBH3 (56 mg, 0.9 mmol) was added and the reaction was allowed to and washed with water (3 x 20 mL) and brine (2x 20 mL). Concentration in vacuo methylaminoethane (114 mg, 0.45 mmol) was added. After 15 minutes at room was then dissolved in ethanol (4 mL, 0.1M) and 1-N, N-dimethylamino-2-N-15

afforded 134 mg (93%) of an orange oil. Column chromatography (9:1, CH₂Cl₂:MeOH) stir overnight at room temperature. The reaction was then with water, diluted with ether afforded an orange oil. LCMS indicated a purity of 99% and hit the mass, 321.2.

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7-OH tetrahydroisoquinoline series

7-Hydroxy-3,4-dihydro-1-H-isoquinoline-2-carboxylic acid tert-butyl ester is prepared by the procedure described in Kucznierz, et.al., J. Med. Chem. 1998, 41, 4983-4994. MS(ES-) 248.1 (M-H):

Example 22

7-(3-Piperidin-1-yl-propoxy-3,4-dihydro-1-H-isoquinoline-2-earboxylic acid tert-butyl

10 ester, Procedure A: A 100 mL dioxane solution of 7-hydro

Procedure A: A 100 mL dioxane solution of 7-hydroxy-3,4-dihydro-1-H-isoquinoline-2-carboxylic acid tear-butyl ester (5.0 g, 20 mmol) is stirred under N₂ as Cs₂CO₃ (13.3 g, 43 mmol), KI (0.1 g, 0.6 mmol), then N-(3-chloropropyl)piperidine (3.9 g, 24 mmol) are added in succession. The reaction mixture is heated at 90°C for 10 hours, cooled, filtered, and concentrated to give the crude product. Purification by chromatography (SiO₂: 0-10% MeOH/CH₂Cl₂/1%NH₄OH gradient) gives the product as an amber oil (7.5 g, 100% yield). MS(ES+)375.3(M+H)*

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Example 238

7-(3-Piperidin-1-yl-propoxy)-1,2,3,4-tetrahydro-isoquinoline dihydrochloride;

Procedure B: A 50 mL CH₂Cl₂ solution of 7-(3-Piperidin-1-yl-propoxy-3,4-dihydro-1-H-isoquinoline-2-carboxylic acid tert-butyl ester (5.1 g, 13.8 mmol) is stirred under N₂ at 0-10⁶C as 4N HCl/dioxane (11.5 mL, 46 mmol) is added dropwise. After the addition is complete, reaction mixture is stirred at this temperature for 30-60 min, then allowed to warm to room temperature. A white precipitate forms and dry MeOH is added until clear solution is obtained. Additional 4N HCl/dioxane (11.0 mL, 44 mmol) is added dropwise.

10 After the addition is complete, reaction mixture is stirred at room temperature. Reaction is followed by TLC (SiO₂ plate, CH₃Cl/McOH/NH₄OH; 25/5/1) until starting material consumed (4-5 h). Reaction mixture is concentrated, dissolved in dry McOH, concentrated, triturated in Et₂O, filtered, and dried in vacuo to give the di-HCl salt (4.5 g, 94% yield) as a white solid. MS(ES+)275.3(M+H)*free base.

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Example 245

2-Methyl-7-(3-Piperidin-1-yl-propoxy)-1,2,3,4-tetrahydro-isoquinoline: A 10 mL THF suspension of LAH (150 mg,4 mmol) is stirred under N₂ at 0-10°C as a 10 mL THF solution of 7-(3-piperidin-1-yl-propoxy-3,4-dihydro-1-H-isoquinoline-2-carboxylic acid

- 20 solution of 7-(3-piperidin-1-yl-propoxy-3,4-dihydro-1-H-isoquinoline-2-carboxylic acid terr-butyl ester (200 mg, 0.53 mmol) is added dropwise. Reaction mixture is allowed to warm to room temperature, refluxed 90 minutes, cooled to 0-10°C, quenched with H₂O and 15% aqueous NaOH, filtered, and the filtrate concentrated to give crude product. Material is purified by chromatography (SiO₂; 0-10% MeOH/CH₂Cl₂/1%NH₄OH
 - 25 gradient)to give the product (82 mg, 54% yld). MS(ES+)289.1(M+H)*.

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Example 2

2-Ethyl-7-(3-Piperidin-1-yl-propoxy)-1.2.3.4-tetrahydro-isoquinoline dihydrochloride; Procedure C; An 80 mL CH,ClyMcOH (9:1) solution of 7-(3-piperidin-1-yl-propoxy)-1,2,3.4-tetrahydro-isoquinoline dihydrochloride (658972)(2.95 g, 8.5mmol) is stirred under N₂, the MP-CNBH₃ resin(15 g, 38 mmol) added, the acetaldehyde (5 mL, 89 mmol) added, the pH is adjusted to -4 with glacial AcOH and reaction mixture stirred at room

temperature for 18-20 hours. The reaction mixture is filtered and the resin beads washed twice alternately with MeOH, then CH₂Cl₂. The filtrate is concentrated and the residue is purified by chromatography (SCX-MeOH wash, elute 2M NH₂/MeOH; then (SiO₂: 0-10% MeOH/CH₂Cl₂/1%NH₂OH gradient) to give the pure free base.

Procedure D: A 50 mL THF/MeOH (1:1) solution of the free base (1.52 g, 5 mmol) is stirred under N₂ at 0-10°C as 1N HCVE₃O (11.5 mL, 11.5 mmol) is added dropwise. After the addition is complete, reaction mixture is allowed to warm to room temperature, then reaction mixture is concentrated, dissolved in dry MeOH, concentrated, triturated in E₂O, filtered, and dried in vacuo to give the di-HCl salt (4.5 g, 94% yld) as a white solid. MS(ES+)303.3(M+H)* free base.

Example 292 (di-HCL salt)

Example 273 (free base)

2-Cyclohexylmethyl-7-(3-piperidin-1-yl-propoxy)-1,2,3,4-tetrahydro-isoquinoline dihydrochloride: 2-Cyclohexylmethyl-7-(3-piperidin-1-yl-propoxy)-1,2,3,4-tetrahydro-isoquinoline is prepared from 7-(3-piperidin-1-yl-propoxy)-1,2,3,4-tetrahydro-isoquinoline dihydrochloride (6 g. 17 mmol), MP-CNBH₃ (30 g. 76.5 mmol), and cyclohexanecarboxaldehyde (12.4 mL, 102 mmol) via a procedure substantially analogous to Procedure C except that the SCX column is not used in purification. The di-

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HCl salt product (4.9 g, 65% yld) is isolated as a white solid via a procedure substantially analogous to Procedure D. MS(ES+)371.4(M+H)*free base.

And alone

2-Isopropyl-7-(3-piperidin-1-yl-propoxy)-1,2,3,4-tetrahydro-isoquinoline: 2-Isopropyl-7-(3-piperidin-1-yl-propoxy)-1,2,3,4-tetrahydro-isoquinoline is prepared from 7-(3-piperidin-1-yl-propoxy)-1,2,3,4-tetrahydro-isoquinoline dihydrochloride (320 mg, 1,5 mmol), MP-CNBH; (3.2 g, 7.5 mmol), and acetone (1.1 mL, 15 mmol) via a procedure substantially analogous to Procedure C except that the SCX column is not used in purification. The product (210 mg, 44% yld) is isolated as a clear oil.

MS(ES+)317.2(M+H)*

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Example 275

15 1-[7-(3-Piperidin-1-yl-propoxy)-3,4-dihydno-1H-isoquinolin-2-yll-ethanone: A 5 mL CH₂Cl₂ solution of 7-(3-piperidin-1-yl-propoxy)-1,2,3,4-tetrahydro-isoquinoline dihydrochloride (175 mg, 0.5 mmol) and NEt₃ (0.25 mL, 1.7 mmol) is stirred under N₂, a 1 mL CH₂Cl₂ solution of acetyl chloride (0.043 mL, 0.6 mmol) is added, and reaction is stirred at room temp. for 5-6 hours. Reaction mixture is quenched with MeOH,

concentrated and the residue is purified by chromatography (SCX-MeOH wash, elute 2M NHyMeOH; then (SiO₃; 0-10% MeOH/CH₃Cl₂1%NH₂OH gradient) to give the product (90 mg, 58% yld). MS(ES+)317.1(M+H)*

[7-(3-Piperidin-1-yl-propoxy)-3,4-dihydro-1H-isoquinolin-2-yl}-thiophen-2-ylmethanone;

propoxy)-1,2,3,4-tetrahydro-isoquinoline dihydrochloride (256 mg, 0.74 mmol), resin bound DCC (1.1 g, 0.9 mmol), hydroxybenzotriazole (HOBt, 150 mg, 1.1 mmol), and Procedure E: A 7 mL CHCl-/r-BuOH/McCN (5:1:1) mixture of 7-(3-piperidin-1-yl-

- twice alternately with MeOH, then CH2Cl2: The filtrate is concentrated and the residue is purified by chromatography (SCX-MeOH wash, elute 2M NHy/MeOH; then SiO2; 0-10% yld). MS(ES+) 385.1(M+H)*. A 3 mL dry McOH solution of the free base (45 mg, 0.12 MeOH/CH₂Cl₂/1% NH₄OH gradient) to give the pure free base as a solid (180 mg, 63% triturated with Et2O, filtered, and dried in vacuo to the HCl salt as an off-white solid (46 mmol) is stirred with 1N HCl/Et₂O (0.18 mL, 0.18 mmol) for 5 minutes, concentrated, temperature for 48 hours. The reaction mixture is filtered and the resin beads washed thiophene-2-carboxylic acid (118 mg, 0.9 mmol) is shaken in a capped vial at room mg). MS(ES+) 385.1(M+H)*free base. 2

Example 274

ethanone: 2-Dimethylamino-1-[7-(3-piperidin-1-yl-propoxy)-3,4-dihydro-1H-isoquinolinresin beads (700 mg, 2.6 mmol) is used in the work up to scavenge the excess HOBt and isoquinoline dihydrochloride (175 mg, 0.5 mmol), PS-DCC (800 mg, 1.1 mmol), HOBt mmol) via a procedure substantially analogous to Procedure E except that PS-trisamine (80 mg, 0.77 mmol), NEt₃ (0.21 mL, 1.5 mmol)and N,N-dimethylglycine (1.1 mL, 15 2-Dimethylamino-1-[7-(3-piperidin-1-yl-propoxy)-3,4-dihydro-1H-isoquinolin-2-yl]-2-yl]-ethanone is prepared from 7-(3-piperidin-1-yl-propoxy)-1,2,3,4-tetrahydro-

53

N.N-dimethylglycine. The free base product (35 mg, 19% yld) is isolated as an oil. MS(ES+)360.5(M+H)*

tetrahydro-isoquinoline dihydrochloride (254 mg, 0.73 mmol), NEts (0.20 mL, 1.4 mmol), concentrated and the residue is purified by chromatography (SCX-MeOH wash, elute 2M NH3/McOH; then SiO₂; 0-10% MeOH/CH₂Cl₂/1%NH,OH gradient) to give pure product mmol) is stirred under N2, at room temperature for 18 hours. The reaction mixture is isopropyl isocyanate (192 mg, 2.2 mmol), and 4-dimethylaminopyridine (12 mg, 0.1 isopropylamide: A 10 mL CH2Cl2 solution of 7-(3-piperidin-1-yl-propoxy)-1,2,3,4-7-(3-Piperidin-1-yl-propoxy)-3,4-dihydro-1H-isoquinoline-2-carboxylic acid (110 mg, 42% yld). MS(ES+) 360.2(M+H) 2

- mmol) is stirred under N2, benzenesulfonyl chloride (0.08 mL, 0.62 mmol) is added, and reaction is stirred at room temperature for 5-6 hours. Reaction mixture is diluted with tetrahydro-isoquinoline dihydrochloride (185 mg, 0.53 mmol) and NEt₃ (0.22 mL, 1.8 2-Benzenesulfonyl-7-(3-piperidin-1-yl-propoxy)-1,2,3,4-tetrahydro-isoquinoline; Procedure F: A 5 mL CH₂Cl₂ solution of 7-(3-piperidin-1-yl-propoxy)-1,2,3,4-15
- residue is purified by chromatography (SiO2; 0-6% MeOH/CH2Cl3/1% NH4OH gradient) with EtOAc. The EtOAc extracts are combined, dried (Na2SO4), and concentrated. The EtOAc, washed with saturated aqueous Na2CO3, and the aqueous layer back-extracted to give the product (160 mg, 73% yld). MS(ES+) 415.1(M+H)*.

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Example 268

7-(3-Piperidin-1-yl-propoxy)-2-(thiophene-2-sulfonyl)-1,2,3,4-tetrahydro-isoquinoline: 7-(3-Piperidin-1-yl-propoxy)-2-(thiophene-2-sulfonyl)-1,2,3,4-tetrahydro-isoquinoline is prepared from 7-(3-piperidin-1-yl-propoxy)-1,2,3,4-tetrahydro-isoquinoline dihydrochloride (175 mg, 0,5 mmol), NEi3 (0.25 mL, 1.8 mmol), and thiophene-2-sulfonyl chloride (114 mg, 0,63 mmol) via a procedure substantially analogous to Procedure F except that an additional SCX column purification step is performed to give

the product (160 mg, 76% yld). MS(ES+)421.1(M+H)

xample 20

7-(3-Piperidin-1-yl-propoxy)-2-(propane-2-sulfonyl)-1,2,3,4-tetrahydro-isoquinoline: 7(3-Piperidin-1-yl-propoxy)-2-(propane-2-sulfonyl)-1,2,3,4-tetrahydro-isoquinoline is prepared from 7-(3-piperidin-1-yl-propoxy)-1,2,3,4-tetrahydro-isoquinoline dihydrochloride (175 mg, 0.5 mmol), NEt₃ (0.25 mL, 1.8 mmol), and isopropylsulfonyl chloride (0.07 mL, 0.60 mmol) via a procedure substantially analogous to Procedure F except that an additional SCX column purification step is performed to give the product (93 mg, 49% yld). MS(ES+) 381.1(M+H)*

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Example 284

- 20 2-Methanesulfonyl-7-(3-piperidin-1-yl-propoxy)-1,2,3,4-tetrahydro-isoquinoline hydrochloride: 2-Methanesulfonyl-7-(3-piperidin-1-yl-propoxy)-1,2,3,4-tetrahydro-isoquinoline hydrochloride is prepared from 7-(3-piperidin-1-yl-propoxy)-1,2,3,4-tetrahydro-isoquinoline dihydrochloride (183 mg, 0.52 mmol), NEty (0.25 mL, 1.8 mmol), and methanelsulfonyl chloride (0.05 mL, 0.66 mmol) via a procedure substantially
 - 25 analogous to Procedure F except that an additional SCX column purification step is performed to give the free base product. A 5 mL dry MeOH solution of the free base (110 mg, 0.31 mmol) is stirred with 1N HC/Et₂O (0.50 mL, 0.5 mmol) for 5 minutes,

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concentrated, triturated with Et₂O, the Et₂O decanted, and the residue dried in vacuo to give the HCl salt as a glass (118 mg, 65% yld). MS(ES+) 353.2(M+H)*free base.

xample 286

- 2-(4-Methoxy-benzenesulfonyl-7-(3-piperidin-1-yl-propoxy)-1,2,3,4-tetrahydro-isoquinoline hydrochloride: 2-(4-Methoxy-benzenesulfonyl-7-(3-piperidin-1-yl-propoxy)-1,2,3,4-tetrahydro-isoquinoline hydrochloride is prepared from 7-(3-piperidin-1-yl-propoxy)-1,2,3,4-tetrahydro-isoquinoline dihydrochloride (150 mg, 0.43 mmol), NEty (0.21 mL, 1.5 mmol), and 4-methoxybenzenesulfonyl chloride (115 mg, 0.57 mmol) via a
 - procedure substantially analogous to Procedure F except that an additional SCX column purification step is performed to give the free base product. A 5 mL dry MeOH solution of the free base (131 mg, 0.29 mmol) is stirred with IN HCUE₁₂O (0.40 mL, 0.4 mmol) for 5 minutes, concentrated, triturated with E₂O, filtered, and dried in vacuo to give the HCl salt (118 mg, 57% yld). MS(ES+) 445.2(M+H)*free base.

Example 277

12

1-{4-[7-(3-Piperidin-1-yl-propoxy)-3,4-dihydro-1H-isoquinoline-2-sulfonyll-phenyl}-ethanone: 1-{4-[7-(3-Piperidin-1-yl-propoxy)-3,4-dihydro-1H-isoquinoline-2-sulfonyl}-phenyl}-ethanone is prepared from 7-(3-piperidin-1-yl-propoxy)-1,2,3,4-tetrahydro-isoquinoline dihydrochloride (175 mg, 0.5 mmol), NEt₃ (0.25 mL, 1.8 mmol), and 4-acetylbenzenelsulfonyl chloride (131 mg, 0.60 mmol) via a procedure substantially analogous to Procedure F except that an additional SCX column purification step is

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performed to give the product (85 mg, 37% yld). MS(ES+) 457.1(M+H)*.

Example 276

tetrahydro-isoquinoline is prepared from 7-(3-piperidin-1-yl-propoxy)-1,2,3,4-tetrahydro isoquinoline dihydrochloride (175 mg, 0.5 mmol), NEi3 (0.25 mL, 1.8 mmol), and 4-(nanalogous to Procedure F except that an additional SCX column purification step is isoquinoline: 2-(4-n-Butyl-benzenesulfonyl)-7-(3-piperidin-1-yl-propoxy)-1,2,3,4butyl)benzenesulfonyl chloride (140 mg, 0.60 mmol) via a procedure substantially 2-(4-n-Butyl-benzenesulfonyl)-7-(3-piperidin-1-yl-propoxy)-1,2,3,4-tetrahydroperformed to give the product (165 mg, 70% yld). MS(ES+)471.1(M+H)*.

Example 278

9

tetrahydro-isoquinoline is prepared from 7-(3-piperidin-1-yl-propoxy)-1,2,3,4-tetrahydroisoquinoline dihydrochloride (175 mg, 0.5 mmol), NEts (0.25 mL, 1.8 mmol), and 4analogous to Procedure F except that an additional SCX column purification step is cyanobenzenesulfonyl chloride (121 mg, 0.60 mmol) via a procedure substantially isoquinoline: 2-(4-Cyanobenzenesulfonyl)-7-(3-piperidin-1-yl-propoxy)-1,2,3,4-2-(4-Cyanobenzenesulfonyl)-7-(3-piperidin-1-yl-propoxy)-1,2,3,4-tetrahydroperformed to give the product (157 mg, 71% yld). MS(ES+) 440.1(M+H)*.

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Example 287

4-[7-(3-Piperidin-1-yl-propoxy)-3,4-dihydro-1H-isoquinoline-2-sulfonyl]- benzamide: A 1.4 mL DMSO mixture of K₂CO₃ is stirred under N₂, 2-(4-cyanobenzenesulfonyl)-7-(3-

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piperidin-1-yl-propoxy)-1,2,3,4-tetrahydro-isoquinoline (75 mg, 0.17 mmol) is added, 0.2 mL H₂O added, followed by 30% H₂O₂(1.4 mL, 12 mmol) and reaction is stirred at room solids washed twice with MeOH. The filtrate is concentrated and the residue is purified temperature for 4 hours. The reaction mixture is diluted with McOH, filtered, and the MeOH/CH₂Cl₂/1%NH₄OH gradient) to give the product as an off-white solid (26 mg, by chromatography (SCX-McOH wash, clute 2M NHy/McOH; then SiO2; 0-10% 26% yld). MS (ES+)458.2(M+H)*.

isoquinoline hydrochloride: 2-(4-Fluoro-benzenesulfonyl)-7-(3-piperidin-1-yl-propoxy)procedure substantially analogous to Procedure F except that an additional SCX column (0.21 mL, 1.5 mmol), and 4-fluorobenzenesulfonyl chloride (115 mg, 0.55 mmol) via a propoxy)-1,2,3,4-tetrahydro-isoquinoline dihydrochloride (158 mg, 0.45 mmol), NEt3 converted to the HCl salt (150 mg, 71% yld) via a procedure substantially analogous purification step is performed to give 140 mg of free base product. The free base is 1,2,3,4-tetrahydro-isoquinoline hydrochloride is prepared from 7-(3-piperidin-1-yl-2-(4-Fluoro-benzenesulfonyl)-7-(3-piperidin-1-yl-propoxy)-1,2,3,4-tetrahydro-Procedure D. MS (ES+)433.2(M+H)*free base.

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Example 304

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tetrahydro-isoquinoline is prepared from 7-(3-piperidin-1-yl-propoxy)-1,2,3,4-tetrahydroisoquinoline dihydrochloride (104 mg, 0.3 mmol), NEt3 (0.14 mL, 1.1 mmol), and 2isoquinoline: 2-(2-Fluoro-benzenesulfonyl)-7-(3-piperidin-1-yl-propoxy)-1,2,3,4fluorobenzenesulfonyl chloride (80 mg, 0.41 mmol) via a procedure substantially 2-(2-Fluoro-benzenesulfonyl)-7-(3-piperidin-1-yl-propoxy)-1,2,3,4-tetrahydro-23

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analogous to Procedure F except that an additional SCX column purification step is performed to give the free base product (85 mg, 66% yld) as an amber oil. MS (ES+) 433.2(M+H)*.

Example 305

2-(3-Fluoro-benzenesulfonyl)-7-(3-piperidin-1-yl-propoxy)-1,2,3,4-tetrahydro-isoquinoline: 2-(3-Fluoro-benzenesulfonyl)-7-(3-piperidin-1-yl-propoxy)-1,2,3,4-tetrahydro-isoquinoline is prepared from 7-(3-piperidin-1-yl-propoxy)-1,2,3,4-tetrahydro-isoquinoline dihydrochloride (104 mg, 0,3 mmol), NEt₃ (0.14 mL, 1.1 mmol), and 3-fluorobenzenesulfonyl chloride (80 mg, 0,41 mmol) via a procedure substantially analogous to Procedure F except that an additional SCX column purification step is performed to give the free base product (90 mg, 70% yld) as an off-white solid. MS

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6-OH tetrahydroisoquinoline series

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6-hydroxy-3,4-dihydro-1H-isoquinoline-2-carboxylic acid tert-butyl ester is prepared by the procedures similar to those described in Selnick, H.G.; Smith, G. R.; Tebben, A. J.; Synth. Commun. 1995, 25, 3255-3262.

Example 127

6-(3-Piperidin-1-yl-propoxy)-3,4-dihydro-1H-isoquinoline-2-carboxylic acid terr-butyl ester: To a round-bottom flask, equipped with stir bar and septum, is placed 6-hydroxy-3,4-dihydro-1H-isoquinoline-2-carboxylic acid tert-butyl ester (1 g. 4.01 mmol), KI (599 mg, 4.01 mmol) and NaH (162 mg, 95%dry, 6.42 mmol). Then, dry DMF (20 mL, 0.5 M) is added via syringe followed by N-(3-chloropropyl)piperidine (0.85 mL, 5.2 mmol). The reaction is allowed to stir at 70 degrees overnight. In the morning, the reaction is quenched with water, extracted into EiOAc (3 x 20 mL) and dried over brine. Column chromatography in 9:1 DCM:MeOH affords 6-(3-piperidin-1-yl-propoxy)-3,4-dihydro-

15 IH-isoquinoline-2-carboxylic acid tert-butyl ester an orange oil (1 g, 67%). Mass sec hit M+1, 375; LCMS >95% @ 230 nm and ELSD.

In a similar manner the Examples 35, 139, and 164 are prepared:

Example 35

6-(3-Dimethylamino-propoxy)-3,4-dihydro-1H-isoquinoline-2-carboxylic acid terr-butyl ester; M+1 335

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6-[3-(2-Methyl-pipendin-1-yl)-propoxy]-3,4-dihydro-1H-isoquinoline-2-carboxylic acid tert-butyl ester; M+1 389

6-(2-Pipendin-1-yl-ethoxy)-3,4-dihydro-1H-isoquinoline-2-carboxylic acid tert-butyl ester, M+1 361.

Example 128

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propoxy)-3,4-dihydro-1H-isoquinoline-2-carboxylic acid tert-butyl ester (1 g, 2.6 mmol), DCM (20 mL) and 4M HCl/dioxane (5 mL). The reaction is allowed to stir at room round-bottom flask, equipped with stir bar and septum, is placed 6-(3-piperidin-1-yl-6-(3-Piperidin-1-yl-propoxy)-1,2,3,4-tetrahydro-isoquinoline dihydrochloride: To a

concentrated again affording 6-(3-piperidin-1-yl-propoxy)-1,2,3,4-tetrahydro-isoquinoline temperature for 3 h. After this time, the reaction is concentrated, dissolved in MeOH and dihydrochloride as a white solid (800 mg, 87%). Mass spec hit M+1, 275; LCMS >95% @ 230 nm and ELSD. 15

In a similar manner the Examples 40, 140, and 165 are prepared:

Dimethyl-[3-(1,2,3,4-tetrahydro-isoquinolin-6-yloxy)-propyl]-amine dihydrochloride; M+1 235.

Example 140

6-[3-(2-Methyl-piperidin-1-yl)-propoxy]-1,2,3,4-tetrahydro-isoquinoline dihydrochloride;

M+1 289.

Example 165

6-(2-Piperidin-1-y1-ethoxy)-1,2,3,4-tetrahydro-isoquinoline dihydrochloride; M+1 261.

Example 129

2

2-Ethyl-6-(3-piperidin-1-yl-propoxy)-1,2,3,4-tetrahydro-isoquinoline: To a 25 mL rounddihydrochloride (700 mg, 2.01 mol), MP-CNBH3 (2.5 g, 6.05 mmol, 2.42 mmol/g) and bottom flask is placed 6-(3-Piperidin-1-yl-propoxy)-1,2,3,4-tetrahydro-isoquinoline DCM/MeOH (9mL/1mL). Then, acetaldehyde is added (0.7 mL, 12 mmol) and the

- DCM/MeOH and concentrated. Column chromatography in 9:1 DCM:MeOH affords 2ethyl-6-(3-piperidin-1-yl-propoxy)-1,2,3,4-tetrahydro-isoquinoline (493 mg, 71%) of a viscous oil. Mass spec hit M+1, 303; LCMS >95% @ 230 nm and ELSD. Атау reaction is allowed to stir overnight. The reaction is then filtered, washed with 15
 - synthesis followed this general procedure in 4 mL vials to make the following 8

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MS	263	320	292	346	326	326	317	329	357	371	329	317	289
Name	[3-(2-Ethyl-1,2,3,4-tetrahydro-isoquinolin-6-yloxy)-propyl]- dimethyl-amine	[3-[6-(3-Dimethylamino-propoxy)-3,4-dihydro-1H-isoquinolin-2-yl]-propyl]-dimethyl-amine	2-[6-(3-Dimethylamino-propoxy)-3.4-dihydro-1H-isoquinolin-2-yl]- acetamide	Dimethyl-(3-[2-(2-piperidin-1-yl-ethyl)-1,2,3,4-tetrahydro-isoquinolin-6-yloxy]-propyl J-amine	Dimethyl-[3-(2-pyridin-3-ylmethyl-1,2,3,4-tetrahydro-isoquinolin-6-yloxyl-propyl]-amine	Dimethyl-[3-(2-pyridin-2-ylmethyl-), 2,3,4-tetrahydro-isoquinolin-6-yloxyl-propyl]-amine	2-Ethyl-6-[3-(2-methyl-piperidin-1-yl)-propoxy]-1,2,3,4-tetrahydro-isoquinoline	2-Cyclopropylmethyl-6-(3-piperidin-1-yl-propoxy)-1,2,3,4-terrahydro-isoquinoline	2-Cyclopentylmethyl-6-(3-piperidin-1-yl-propoxy)-1,2,3,4-terrahydro-isoquinoline	2-Cyclohexylmethyl-6-(3-piperidin-1-yl-propoxy)-1,2,3,4-tetrahydro-isoquinoline	2-(2-Ethyl-butyl)-6-(3-piperidin-1-yl-propoxy)-1,2,3,4-tetrahydro-isoquinoline	6-(3-Piperidin-1-yl-propoxy)-2-propyl-1,2,3,4-tetrahydro- isoquinoline	2-Ethyl-6-(2-piperidin-1-yl-ethoxy)-1,2,3,4-tetrahydro-isoquinoline
Example	92	11	08	81	82	83	141	145	146	147	148	149	166

345 33 2-Cyclopropylmethyl-6-(2-piperidin-1-yl-ethoxy)-1,2,3,4-tetrahydro- 315 2-Cyclohexylmethyl-6-(2-piperidin-1-yl-ethoxy)-1,2,3,4-tetrahydro- 357 2-Cyclopentylmethyl-6-(2-piperidin-1-yl-ethoxy)-1,2,3,4-tetrahydro-2-(2-Ethyl-butyl)-6-(2-piperidin-1-yl-ethoxy)-1,2,3,4-tetrahydro-2-Isopropyl-6-(2-piperidin-1-yl-ethoxy)-1,2,3,4-tetrahydroisoquinoline isoquinoline isoquinoline isoquinoline isoquinoline 17 168 172

Example 250

MeOH (50 mL), and 1M HCl in ether is added dropwise (37.2 mL, 37.2 mmol) and the mixture is 5 2-Ethyl-6-(3-piperidin-1-yl-propoxy)-1,2,3,4-tetrahydro-isoquinoline dihydrochloride: 2-Ethyl-6stirred for 10 minutes and concentrated to give the dihydrochloride salt as a white solid (6.0 g. (3-piperidin-1-yl-propoxy)-1,2,3,4-tetrahydro-isoquinoline (5.12g, 16.9 mmol) is dissolved in 93%).

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Example 143

a flask equipped with a stir bar is placed 6-[3-(2-Methyl-piperidin-1-yl)-propoxy]-1,2,3,4-2-Isopropyl-6-[3-(2-methyl-piperidin-1-yl)-propoxy]-1,2,3,4-tetrahydro-isoquinoline: To tetrahydro-isoquinoline dihydrochloride (300 mg, 0.83 mmol), acetone (excess),

temperature for 2h. The reaction mixture is diluted with water, and extracted with NaCNBH₃ (155 mg, 2.5 mmol) in MeOH (8 mL) and the mixture stirred at room 2

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CH2Cl2. The organic phase is dried over Na₂SO₄ and concentrated. M+1 331, LCMS >98% @ 230 nm and ELSD.

In a similar manner Example 138 is prepared:

Example 138

2-Isopropyl-6-(3-piperidin-1-yl-propoxy)-1,2,3,4-tetrahydro-isoquinoline; M+1 317, LCMS 100% @ 230 nm and ELSD.

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added and the reaction is again allowed to rotate overnight to scavenge excess carboxylic [6-(3-Piperidin-1-yl-propoxy)-3,4-dihydro-1H-isoquinolin-2-yl]-thiazol-2-yl-methanone: mmol/g), HOBt (16 mg, 0.12 mmol), pyrazole carboxylic acid (13 mg, 0.1 mmol) and a pipendin-1-yl-propoxy)-3,4-dihydro-1H-isoquinolin-2-yl]-thiazol-2-yl-methanone as an synthesis follows this general procedure in 4 mL vials to make the following examples: orange foam. Filtration through a short pipet column provides 24 mg (80%) of [6-(3-To a 4 mL vial is placed 6-(3-piperidin-1-y)-propoxy)-1,2,3,4-tetrahydro-isoquinoline 5:1:1 mixture of CHCl₃:CH₃CN:tBuOH. The vial is agitated by means of a lab quake shaker overnight. In the morning, PS-trisamine (134 mg, 0.4 mmol, 3.0 mmol/g) is acid and HOBt. Filtration, washing with DCM/MeOH and concentration affords a orange solid. Mass spec hit M+1, 386; LCMS >95% @ 230 nm and ELSD. Аттау dihydrochloride (28 mg, 0.08 mmol), resin-bound DCC (134 mg, 0.16 mmol, 1.2

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zample	Name	MS
78	[6-(3-Dimethylamino-propoxy)-3,4-dihydro-1H-isoquinolin-2-yl]-	474
	(1-phenyl-5-trifluoromethyl-1H-pyrazol-4-yl)-methanone	

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315	386	383	368	363	385	402	386	386	360	386	346	332	344	358
1-[6-(3-Piperidin-1-yl-propoxy)-3,4-dihydro-1H-isoquinolin-2-yl]- ethanone	[6-(3-Piperidin-1-yl-propoxy)-3,4-dihydro-1H-isoquinolin-2-yl]- (tetrahydro-furan-2-yl)-methanone	(5-Methyl-furan-2-yl)-[6-(3-piperidin-1-yl-propoxy)-3,4-dihydro- IH-isoquinolin-2-yl]-methanone	[6-(3-Piperidin-1-yl-propoxy)-3,4-dihydro-1H-isoquinolin-2-yl]- (1H-pyrrol-2-yl)-methanone	2-Methylsulfanyl-1-[6-(3-piperidin-1-yl-propoxy)-3,4-dihydro-1H-isoquinolin-2-yl]-ethanone	[6-(3-Piperidin-1-yl-propoxy)-3,4-dihydro-1H-isoquinolin-2-yl]- thiophen-2-yl-methanone	N.N.Dimethyl-4-oxo-4-[6-(3-piperidin-1-yl-propoxy)-3,4-dihydro- 1H-isoquinolin-2-yl]-butyramide	[6-(3-Piperidin-1-yl-propoxy)-3,4-dihydro-1H-isoquinolin-2-yl]-thiazol-2-yl-methanone	5-[6-(3-Piperidin-1-yl-propoxy)-3,4-dihydro-1H-isoquinoline-2-carbonyl]-pyrrolidin-2-one	2-Dimethylamino-1-{6-(3-piperidin-1-yl-propoxy)-3,4-dihydro-1H-isoquinolin-2-yl]-ethanone	(1-Methyl-pyrrolidin-2-yl)-[6-(3-piperidin-1-yl-propoxy)-3,4-dihydro-1H-isoquinolin-2-yl]-methanone	2-Dimethylamino-1-[6-(2-piperidin-1-yl-ethoxy)-3,4-dihydro-1H-isoquinolin-2-yl]-ethanone	1-[6-(3-Piperidin-1-yl-propoxy)-3.4-dihydro-1H-isoquinolin-2-yl]- propan-1-one	Cyclopropyl-[6-(3-piperidin-1-yl-propoxy)-3,4-dihydro-1H-isoquinolin-2-yl]-methanone	Cyclobutyl-[6-(3-piperidin-1-yl-propoxy)-3,4-dihydro-1H-isoquinolin-2-yl]-methanone
134	136	157	158	128	91	191	162	163	271	176	171	182	183	184

In a similar manner Examples 179 is prepared:

346

2-Methyl-1-[6-(3-piperidin-1-yl-propoxy)-3,4-dihydro-1H-

186

Cyclopentyl-[6-(3-piperidin-1-yl-propoxy)-3,4-dihydro-1H-

isoquinolin-2-yl]-methanone

385

Cyclohexyl-[6-(3-piperidin-1-yl-propoxy)-3,4-dihydro-1H-

187

isoquinolin-2-yl]-propan-1-one

2-Ethyl-1-[6-(3-piperidin-1-yl-propoxy)-3,4-dihydro-1H-

188

isoquinolin-2-yl]-methanone

isoquinolin-2-yl]-butan-1-one

Example 179

[3-(2-Methanesulfonyl-1,2,3,4-tetrahydro-isoquinolin-6-yloxy)-propyl]-dimethyl-amine: (12 DL, 0.15 mmol) and dry CH₂Cl₂ (2 mL). The vial is allowed to rotate overnight. In To a 4 mL vial is placed Dimethyl-[3-(1,2,3,4-tetrahydro-isoquinolin-6-yloxy)-propyl]amine (24.0 mg, 0.1 mmol), resin-bound DIEA (58 mg, 0.2 mmol, 3.54 mmol/g), MsCl the morning, PS-trisamine (136 mg, 0.41 mmol, 3.0 mmol/g) is added and the reaction 2

CH₂Cl₂ and concentration affords the desired urea LCMS >99% @ 230 nm and ELSD, again allowed to rotate for 4 hours to scavenge excess MsCl. Filtration, washing with 12

Example 302

tetrahydro-isoquinoline dihydrochloride (25.0 mg, 0.07 mmol), resin-bound Hunigs base

isopropylamide: To a 4 mL vial is placed 6-(3-piperidin-1-yl-propoxy)-1,2,3,4-6-(2-Piperidin-1-yl-propoxy)-3,4-dihydro-1H-isoquinoline-2-carboxylic acid

Example 178

isopropyl isocyanaté (16 IIL, 0.18 mmol). The vial is agitated by means of a lab quake (81 mg, 0.29 mmol, 3.54 mmol/g), resin bound DMAP (catalytic), and dry CH₂Cl₂ and

2

shaker overnight. In the morning, PS-trisamine (120 mg, 0.36 mmol, 3.0 mmol/g) is

added and the reaction again allowed to rotate for 4 hours to scavenge excess isocyanate. Filtration, washing with CH2Cl2 and concentration afforded the desired urea. M+1 360.

Benzenesulfonyl-6-(3-piperidin-1-yl-propoxy)-1,2,3,4-tetrahydro-isoquinoline is prepared mg, 0.95 mmol), NEt₃ (0.48 mL, 3.5 mmol), and benzenesulfonyl chloride (0.15 mL, 1.17 from 6-(3-piperidin-1-yl-propoxy)-1,2,3,4-tetrahydro-isoquinoline dihydrochloride (330 2-Benzenesulfonyl-6-(3-piperidin-1-yl-propoxy)-1,2,3,4-tetrahydro-isoquinoline: 2-ន

6-(2-Pipendin-1-yl-propoxy)-3,4-dihydro-1H-isoquinoline-2-carboxylic acid

cyclohexylamide; M+1 400.

[6-(3-Piperidin-1-yi-propoxy)-3,4-dihydro-1H-isoquinolin-2-yi]-

pyridin-3-yl-methanone

Isoxazol-5-yl-[6-(3-piperidin-1-yl-propoxy)-3,4-dihydro-1H-

196

pyridin-2-yl-methanone

isoquinolin-2-yl]-methanone

[6-(3-Piperidin-1-yl-propoxy)-3,4-dihydro-1H-isoquinolin-2-yl]-

193

[6-(3-Piperidin-1-yl-propoxy)-3,4-dihydro-1H-isoquinolin-2-yl]-

194

pyridin-4-yl-methanone

33

SCX column purification step is performed to give the product as a white solid (250 mg, mmol) via a procedure substantially analogous to Procedure Fexcept that an additional 63% yld). MS(ES+) 415.3(M+H)*.

5-OH tetrahydroisoquinoline series

5-Hydroxy-3,4-dihydro-1-H-isoquinoline-2-carboxylic acid tert-butyl ester is prepared by the procedures similar to those described in Durand S.; Lusinchi, X.; Moreau, R. C. Bull. Soc. Chim. France 1961, 207, 270, and Georgian, V.; Harrison, R. J.; Skaletzky, L. L.; J Org Chem 1962, 27, 4571.

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Example 290

5-(3-Piperidin-1-yl-propoxy)-3,4-dihydro-1H-isoquinoline-2-carboxylic acid tert-butyl ester is prepared from 5-Hydroxy-3,4-dihydro-1-H-isoquinoline-2-carboxylic acid tertbutyl ester (5.69 g, 22.8 mmol) in a manner substantially analogous to Procedure A

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except DMF is used in place of dioxane. Following aqueous workup, the crude material CHClyMeOH/NH,OH) / 90% (10% McOH/CHCl₃)] to give the title compound (5.2 g, is purified by flash chromatography [Biotage 65M SiO₂, elute 10% (25/5/1 61%). MS (ES+) 375.3

Example 291

acid tert-butyl ester (4.0 g, 10.7 mmol) in a manner substantially analogous to Procedure prepared from 5-(3-Piperidin-1-yl-propoxy)-3,4-dihydro-1H-isoquinoline-2-carboxylic 5-(3-Piperidin-1-yl-propoxy)-1,2,3,4-tetrahydro-isoquinoline dihydrochloride salt is B to give the title compound as an off-white solid (3.47 g, 93%). MS (ES+) 275.2

2

Example 309

methanone is prepared from 5-(3-Piperidin-1-yl-propoxy)-1,2,3,4-tetrahydro-isoquinoline Procedure E to give the title compound as an off-white solid (0.109 g, 38%). MS (ES+) dihydrochlonde salt (0.256 g, 0.74 mmol) in a manner substantially analogous to [5-(3-Piperidin-1-yl-propoxy)-3,4-dihydro-1H-isoquinolin-2-yl]-thiophen-2-yl-

21

415.2

Example 294

2

dihydrochloride salt (150 mg, 0.43 mmol) via a procedure substantially analogous to 2-Benzenesulfonyl-5-(3-piperidin-1-yl-propoxy)-1,2,3,4-tetrahydro-isoquinoline is prepared from 5-(3-Piperidin-1-yl-propoxy)-1,2,3,4-tetrahydro-isoquinoline

35

Procedure F to provide the title compound as an off-white solid (54 mg, 30%). MS (ES+) 385.2

Example 306

2-Ethyl-5-(3-pipendin-1-yl-propoxy)-1,2,3,4-tetrahydro-isoquinoline is prepared from 5-(3-Piperidin-1-yl-propoxy)-1,2,3,4-tetrahydro-isoquinoline dihydrochloride salt (375 mg. 1.1 mmol) in a manner substantially analogous to Procedure C to give the title compound as a yellow oil (49 mg, 15%). MS (ES+) 303.3

Example 313

2-Cyclohexylmethyl-5-(3-piperidin-1-yl-propoxy)-1,2,3,4-tetrahydro-isoquinoline is Procedure C to give the title compound as a yellow oil (0.142 mg, 38%). MS (ES+) dihydrochloride salt (350 mg, 1.0 mmol) in a manner substantially analogous to prepared from 5-(3-piperidin-1-yl-propoxy)-1,2,3,4-tetrahydro-isoquinoline 371.4

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8-OH tetrahydroisoquinoline series

8-Methoxy-1,2,3,4-tetrahydro-isoquinoline is prepared according to Shanker, P. S.; Subba Rao, G. S. R. Indian J. of Chemistry section B 1993, 32B, 1209-1213.

78 °C is added a solution of boron tribromide in CH₂Cl₂ (1 M, 52 mL, 52 mmol) dropwise of 8-methoxy-1,2,3,4-tetrahydro-isoquinoline (2.54 g, 15.6 mmol) in CH2Cl2 (60 mL) at over approximately 20 minutes. The cooling bath is removed, and the mixture is warmed 8-Hydroxy-3,4-dihydro-1H-isoquinoline-2-carboxylic acid tert-butyl ester: To a mixture to room temperature. After 4 h, the reaction is carefully quenched with ice. EtOAc and water is added, and the mixture is stirred overnight. The phases are separated, and 5 N 2

room temperature overnight. EtOAc is added, and the phases are separated. The aqueous

phase is extracted with EtOAc (1X), and the combined organic phase is washed with

di-ten-butyl dicarbonate (6.78 g, 31 mmol) is added, and reaction mixture is stirred at

NaOH solution is added to the aqueous phase until pH is basic. Dioxane (250 mL) and

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brine and dried (MgSO₄). After filtration, the solvent is removed in vacuo to provide the title compound (4.84 g) that is used without purification. MS (ES-) 248.2.

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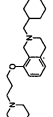
8-(3-Piperidin-1-yl-propoxy)-3,4-dihydro-1H-isoquinoline-2-carboxylic acid tert-butyl ester is prepared from 8-hydroxy-3,4-dihydro-1H-isoquinoline-2-carboxylic acid tert-butyl ester (0.84 g, 3.4 mmol) in a manner substantially analogous to Procedure A except DMF is used in place of dioxane. Following aqueous workup, the crude material is purified by chromatography [SCX-McOH wash, clute 2M NHyMcOH then Biotage 40s SiO₂, elute 10% (25/5/1 CHClyMcOH/NH₂OH) / 90% (10% McOH/CHCl₃)] to give the

title compound (0.61 g, 48%). MS (ES+) 375.3.

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Example 308

8-(3-Pipendin-1-yl-propoxy)-1,2,3,4-terrahydro-isoquinoline dihydrochlonde salt is prepared from 8-(3-pipendin-1-yl-propoxy)-3,4-dihydro-1H-isoquinoline-2-carboxylic acid ten-butyl ester (3.09 g. 8.25 mmol) in a manner substantially analogous to Procedure B to give the title compound as an off-white solid (2.63 g. 85%). MS (ES+) 275.3



Example 309

2-Cyclohexylmethyl-8-(3-piperidin-1-yl-propoxy)-1,2,3,4-tetrahydro-isoquinoline is prepared from 8-(3-piperidin-1-yl-propoxy)-1,2,3,4-tetrahydro-isoquinoline

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dihydrochloride salt (0.375 g, 1.1 mmol) in a manner substantially analogous to Procedure C to give the title compound as a yellow oil (0.195 g, 48%). MS (ES+) 371.4

Example 310

2-Ethyl-8-(3-piperidin-1-yl-propoxy)-1,2,3,4-tetrahydro-isoquinoline is prepared from 8-(3-piperidin-1-yl-propoxy)-1,2,3,4-tetrahydro-isoquinoline dihydrochloride salt (0.375 g, 1.1 mmol) in a manner substantially analogous to Procedure C to give the title compound as a yellow oil (0.124 g, 37%). MS (ES+) 303.3.

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Example 311

2-Benzenesulfonyl-8-(3-piperidin-1-yl-propoxy)-1,2,3,4-tetrahydro-isoquinoline is prepared from 8-(3-piperidin-1-yl-propoxy)-1,2,3,4-tetrahydro-isoquinoline dihydrochloride salt (300 mg, 0.86 mmol) via a procedure substantially analogous to Procedure F to provide the title compound as an off-white solid (0.22 g, 63%). MS (ES+)

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Example 312

[8-(3-Piperidin-1-yl-propoxy)-3,4-dihydro-1H-isoquinolin-2-yl-thiophen-2-yl-

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methanone: To a mixture of 8-(3-pipendin-1-yl-propoxy)-1,2,3,4-tetrahydro-isoquinoline dihydrochloride salt (300 mg, 0.86 mmol) and NE₁₃ (0.36 mL, 2.6 mmol) in CH₂Cl₂ (10 mL) is added 2-thiophene carbonyl chloride (0.10 mL, 0.95 mmol). After stirring at room temperature overnight, the mixture is partitioned between EtOAc and water. The organic phase is washed with brine, dried (MgSO₄), and concentrated. The residue is purified by

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flash chromatography [Biotage 40S SiO₃, elute 20% (90/10/1 CH₂Cl₂/MeOH/NH₄OH) / 80% CH₂Cl₂ to 100% (90/10/1 CH₂Cl₂/MeOH/NH₄OH)] to yield the title compound as a yellow oil (0.181 g, 55%). MS (ES+) 385.3.

Example 206

6-(3-Pipendin-1-yl-propoxy)-3,4-dihydro-2H-isoquinolin-1-one is prepared from 6-hydroxy-3,4-dihydro-2H-isoquinolin-1-one (CAS Registry Number 22245-98-3) (0.5 g, 2.9 mmol) in a manner substantially analogous to Procedure A except DMF is used in place of dioxane. Following aqueous workup, the crude material is purified by flash chromatography (Biotage 40M SiO₂, elute 90/10/1 CH₂Cl₂/MeOH/NH₄OH) to give the title compound as a white solid (0.516 g, 61%). MS (ES+) 289.1

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Example 207

7-(3-Piperidin-1-yl-propoxy)-3,4-dihydro-2H-isoquinolin-1-one is prepared from 7-hydroxy-3,4-dihydro-2H-isoquinolin-1-one (CAS Registry Number 22246-05-5) (1.43 g, 8.76 mmol) in a manner substantially analogous to Procedure A except DMF is used in place of dioxane. Following aqueous workup, the crude material is purified by flash chromatography (Biotage 40M SiO₂, elute 90/10/1 CH₂Cl₃/MeOH/NH₄OH) to give the title compound as a white solid (1.11 g, 44%). MS (ES+) 289.1

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OC alone

7-(3-Pyrrolidin-1-yl-propoxy)-3,4-dihydro-2H-isoquinolin-1-one is prepared from 7-hydroxy-3,4-dihydro-2H-isoquinolin-1-one (0.48 g, 2.94 mmol) in a manner substantially analogous to Procedure A except DMF is used in place of dioxane and 1-(3-Chloropropyl)-pyrrolidine is used instead of N-(3-chloropropyl)piperidine. Following aqueous workup, the crude material is purified by flash chromatography (Biotage 40M SiO₂, elute 90/10/1 CH₂Cl₂/MeOH/NH₄OH) to give the title compound as an off-white solid (0.17 g, 21%). MS (BS+) 275.1

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2-Ethyl-6-hydroxy-3,4-dihydro-2H-isoquinolin-1-one:

To a mixture of 6-methoxy-3,4-dihydro-2H-isoquinolin-1-one (0.30 g, 1.69 mmol) in THF (10 mL) is added sodium hydride (60% mineral oil suspension, 100 mg). The suspension is heated at reflux for 1 h, and cooled to room temperature. Ethyl iodide (1.4 mL, 17 mmol) is added, and the mixture is stirred at room temperature overnight. The mixture is partitioned between EtOAc and water. After the aqueous phase is extracted with EtOAc (2x), the combined organic phase is washed with brine and dried (MgSO₄). After removal of the solvent, the residue is purified by flash chromatography (Biotage After removal of the solvent, the residue is purified by flash chromatography (Biotage After removal of the solvent, the residue is purified by flash chromatography (Biotage After removal of the solvent, the residue is purified by flash chromatography (Biotage After removal of the solvent, the residue is purified by flash chromatography (Biotage After removal of the solvent, the residue is purified by flash chromatography (Biotage After removal of the solvent, the residue is purified by flash chromatography (Biotage After removal of the solvent, the residue is purified by flash chromatography (Biotage After removal of the solvent, the residue is purified by flash chromatography (Biotage After removal of the solvent, the residue is purified by flash chromatography (Biotage After removal of the solvent, the residue is purified by flash chromatography (Biotage After removal of the solvent, the residue is purified by flash chromatography (Biotage After removal of the solvent the solvent

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40M SiO₂, elute 45% EtOAc:hexane – 50% EtOAc:hexane, linear gradient) to yield 2-ethyl-6-methoxy-3,4-dihydro-2H-isoquinolin-1-one as a colorless oil (0.275 g, 78%).
 The material is dissolved in CH₂Cl₂ (10 mL) and cooled to -78 °C. To the cooled mixture is added a solution of boron tribromide (1 M, 4.7 mL, 4.7 mmol) in CH₂Cl₂. After 0.5 h, the temperature is warmed to 0 °C and stirred for 3 h. After the reaction is carefully quenched with ice, EtOAc and water is added, and the mixture is vigorously stirred overnight. The phases are separated, and the organic phase is extracted with EtOAc (2x). The combined organic phase is washed with brine and dried (MgSO₄). The solvent is

removed in vacuo, and the residue is purified by chromatography (Varian 10 g SiO₂

23

cartridge, elute 60% EtOAc:hexane) to provide 2-ethyl-6-hydroxy-3.4-dihydro-2H-isoquinolin-1-one (0.209 g, 82%). MS (ES+) 192.0

xample 265

2-Ethyl-6-(3-piperidin-1-yl-propoxy)-3,4-dihydro-2H-isoquinolin-1-one is prepared from
 2-Ethyl-6-hydroxy-3,4-dihydro-2H-isoquinolin-1-one (0.192 g, 1.0 mmol) in a manner substantially analogous to Procedure A except DMF is used in place of dioxane.
 Following aqueous workup, the crude material is purified by chromatography (Varian 10 g SiO₂ cartridge, elute 10% (25/5/1 CHCI3/MeOH/NH₄OH) / 90% (10% MeOH/CHCI₃)
 to obtain the title compound as a waxy off-white solid (77 mg, 24%). MS (ES+) 317.1

Example 303

[3-Fluoro-4-(3-piperidin-1-yl-propoxy)-phenyl]-(2-pyrrolidin-1-ylmethyl-pyrrolidin-1-yl)-methanone:

General Procedure G: A mixture of (3-Fluoro-4-hydroxy-phenyl)-(2-pytrolidin-1-ylmethyl-pytrolidin-1-yl)-methanone (0.193 g, 0.66 mmol), Cs₂CO₃ (0.43 g, 1.32 mmol), KI (55 mg, 0.33 mmol), and N-(3-chloropropyl)piperidine (3.9 g, 24 mmol) in DMF (5 mL) is heated at 90 °C overnight. The mixture is partitioned between EtOAc and water. The phases are separated, and the aqueous phase is extracted with EtOAc (2x). The combined organic phase is washed with brine, dried (MgSO₄), and concentrated in vacuo. The residue is purified by chromatography (SCX-MeOH wash, elute 2M NH₂/MeOH; then Biotage 12M SiO₂, elute 10% (25/5/1 CHCl₂/MeOH/NH₂OH) / 90% (10% MeOH/CHCl₃)] to give the title compound as a yellow oil (0.105 g, 38%). MS (ES+)

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Example 240

[1-[4-(3-Piperidin-1-yl-propoxy)-phenyl]-cyclopropyl]-carbamic acid benzyl ester is prepared from [1-(4-Hydroxy-phenyl)-cyclopropyl]-carbamic acid benzyl ester (1.21 g.

4.28 mmol), Cs₂CO₃ (2.78 g, 8.55 mmol), KI (71 mg, 0.43 mmol), and N-(3-chloropropyl)piperidine (0.86 g, 5.34 mmol) in dioxane (50 mL) in a manner substantially analogous to Procedure A to give the product (1.16 g, 66%). MS (ES+) 409.3.

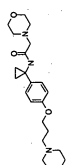
Example 241

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1-[4-(3-Piperidin-1-yl-propoxy)-phenyl]-cyclopropylamine:

[1-[4-(3-Piperidin-1-yl-propoxy)-phenyl]-cyclopropyl]-carbamic acid benzyl ester (1.08 g, 2.65 mmol) is dissolved in ethanol (50 mL), and 10% Pd/C is added (200 mg). The mixture was stirred under a balloon on hydrogen for 3 hours. The reaction mixture was

15 stirred through a plug of silica gel to give the desired compound. HRMS 275.2123 (M+H)*.



Example 247

2-Morpholin-4-yl-N-(1-[4-(3-piperidin-1-yl-propoxy)-phenyl]-cyclopropyl}-acetamide: 1-[4-(3-Piperidin-1-yl-propoxy)-phenyl]-cyclopropylamine (0.195 g, 0.72 mmol) and

NHyMcOH; then Biotage 12M SiO2, elute 10% (25/5/1 CHCly/MeOH/NH,OH) / 90% temperature. The residue is purified by chromatography [SCX-McOH wash, clute 2M diisopropylethylamine added (0.15 mL), followed by EDC (0.165 g, 0.86 mmol) and (10% McOH/CHCl3)) to give the title compound as a yellow oil. HRMS 402.2765 HOBi (0.116 g, 0.86 mmol). The reaction mixture was stirred overnight at room morpholin-4-yl-acetic acid (0.125 g, 0.86 mmol) are dissolved in DMF, and (M+H)

Example 316

2

carboxylic acid tert-butyl ester(1.0 g, 3 mmol), piperidine (0.75 mL, 7.5 mmol), and KI (1.0 g, 6 mmol) is stirred at 50 °C under N2 for four hours, then at room temperature for ester. A 20 mL DMF mixture of 7-(4-chloro-butoxy)-3,4-dihydro-1-H-isoquinoline-2-7-(4-Piperidin-1-yl-butoxy)-3,4-dihydro-1H-isoquinoline-2-carboxylic acid tert-butyl wash, clute 2M NH3/McOH, then SiO2; 0-6% McOH/CH2Cl3/1%NHOH gradient)to I6 hours. The reaction mixture is directly purified by chromatography (SCX-MeOH give the free base (700 mg, 60% yld). MS(ES+)389.3 (M+H)*free base.

2

Example 314

Piperidin-1-yl-butoxy)-1,2,3,4-tetrahydro-isoquinoline dihydrochloride is prepared from 7-(4-chloro-butoxy)-3,4-dihydro-1-H-isoquinoline-2-carboxylic acid tert-butyl ester(600 mg, 1.5 mmol) and 4N HCV dioxane (2.5 mL, 10 mmol) base in a manner substantially 7-(4-Piperidin-1-yl-butoxy)-1,2,3,4-tetrahydro-isoquinoline dihydrochloride: 7-(4analogous to Procedure B to give the product(490 mg, 90% yld). MS(ES+)389.3 (M+H)*free 23

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2-Ethyl-7-(4-piperidin-1-yl-butoxy)-1,2,3,4-tetrahydro-isoquinoline dihydrochloride: 2dihydrochloride (252 mg, 0.7 mmol), and acetaldehyde (0.40 mL, 7 mmol) in a manner Ethyl-7-(4-piperidin-1-yl-butoxy)-1,2,3,4-tetrahydro-isoquinoline dihydrochloride is substantially analogous to Procedure C to give the dihydrochloride product as an off prepared from 7-(4-piperidin-1-yl-butoxy)-1,2,3,4-tetrahydro-isoquinoline white solid(125 mg, 70% yld). MS(ES+)317.2(M+H)* free base.

cyclohexanecarboxaldehyde (0.35 mL, 2.9 mmol) in a manner substantially analogous to Procedure C to give the dihydrochloride product as an off white solid(105 mg, 62% yld). dihydrochloride: 2-Cyclohex ylmethyl-7-(4-piperidin-1-yl-butoxy)-1,2,3,4-tetrahydroisoquinoline dihydrochloride is prepared from 7-(4-piperidin-1-yl-butoxy)-1,2,3,4-2-Cyclohexylmethyl-7-(4-pipendin-1-yl-butoxy)-1,2,3,4-tetrahydro-isoquinoline tetrahydro-isoquinoline dihydrochloride (175 mg, 0.48 mmol), and MS(ES+)385.3(M+H)* free base. 2

Example 208

amination is run with 3-(3-piperidin-1-yl-propoxy)-benzaldehyde (1 g, 4 mmol) and), 3-[3-(3-Piperidin-1-yl-propoxy)-benzyl]-(3-pyrrolidin-1-yl-propyl)-amine: The reductive 23

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pyrrolridin-1-yl propylamine (1 m<u>I.</u>, 8 mmol), and MP-CNBH3 resin(4.5g, 10.4 mmol)via a procedure substantially analogous to [2-(3-piperidin-1-yl-propoxy)-benzyl]-(3-pyrrolidin-1-yl-propyl)-amine to give the product as a yellow oil(818 mg, 58 % yld). MS(ES+)360.3(M+H)* free base.

[4-(4-Pipendin-1-yl-butoxy)-benzyl]-(2-pyrrolidin-1-yl-ethyl)-amine: An 8 mL DMF solution of 14-(4-hromo-hutoxyl-benzyl]-(2-pyrrolidin-1-yl-ethyl)-amine (307 mg, 0.86

- solution of [44(4-bromo-butoxy)-benzyl]-(2-pyrrolidin-1-yl-ethyl)-amine (307 mg, 0.86 mmol) and piperidine (0.22 mL, 2.2 mmol) is stirred at 90 °C for six hours under N₂. The reaction mixture is cooled, diluted with CH₂Cl₂, filtered, washed with brine, dried (Na₂SO₄), and concentrated. The residue is purified by chromatography (SiO₂: 0-6% McOH/CH₂Cl₂/1%NH₄OH gradient) to give the product (40 mg, 12% yld).
- 15 MS(ES+)360.4(M+H)* free base.

Example 236

N-(2-Pipenidin-1-yl-ethyl)-4-(3-pipenidin-1-yl-propox y)-benzamide is prepared according to general procedure A from 4-Hydroxy-N-(2-pipenidin-1-yl-ethyl)-benzamide (CAS Registry 106018-38-6) (0.27 g. 1.1 mmol) to give the title compound as a white solid (77 mg. 19%). MS (ES+) 374.3

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Example 237

2-Fluoro-N-(2-piperidin-1-yl-ethyl)-4-(3-piperidin-1-yl-propoxy)-benzamide:

To a mixture of 2-Fluoro-4-(3-piperidin-1-yl-propoxy)-benzoic acid (70 mg, 0.25 mmol) and 1-(2-aminoethyl)piperidine (45 GL, 0.3 mmol) in DMF (5 mL) was added EDC (58 mg, 0.3 mmol). HOBT (40 mg, 0.3 mmol), and diisopropylethyl amine (52 Gl, 0.3 mmol). The mixture was stirred at room temperature overnight. The mixture was partitioned between EtOAc and water. The organic phase was washed with brine, dried (MgSO₄), and concentrated. The residue was purified by flash chromatography (Biotage 12 M, clute 90/10/1 CH₂Cl₂/MeOH/NH₄OH) to yield the title compound. MS (ES+)

Example 264

3-Fluoro-N-(2-pipendin-1-yl-ethyl)-4-(3-pipendin-1-yl-propoxy)-benzamide is prepared from 3-Fluoro-4-hydroxy-N-(2-pipendin-1-yl-ethyl)-benzamide (0.1 g, 0.38 mmol) by general procedure A to yield the title compound as an off-white solid (80 mg, 54%). MS (ES+) 392.2

12

xample 256

(2-Morpholin-4-yl-ethyl)-[4-(3-piperidin-1-yl-propoxy)-benzyl]-amine dihydrochloride: The dihydrochloride salt was prepared from (2-morpholin-4-yl-ethyl)-[4-(3-piperidin-1-yl-propoxy)-benzyl]-amine (0.307 g) by dissolving in THF (6 mL) and adding a solution

of HCl in Et₂O (1 M, 0.85 mL). Additional Et₂O was added until the mixture was cloudy, and the mixture was allowed to stand at 0 °C overnight. The white solid was collected by filtration to give the dihydrochloride salt. Anal. Calculated for C21H35N3O2 2 HCl: C, 58.06; H, 8.58; N, 9.67; Cl, 16.32. Found: C, 58.0; H, 8.51; N, 9.57; Cl, 16.99.

Synthesis of (1)

- Dimethylglycine (20.0mmol) and 2.58g of N, N-Di-isopropylethylamine (20.0mmol) were dissolved in 50ml of CH₂Cl₂ and 6.78g of PyBOP (13.0mmol) was added to the mixture. The reaction mixture was stirred at room temperature for 4h. The reaction mixture was diluted with 20ml of CH2Cl2 and washed with brine, 0.1N Hl, brine 1.50g of @(+)-1-(4-methoxyphenyl) ethylamine (10.0mmol), 2.06g of N, N-
- $(CH_2Cl_2 \rightarrow CH_2Cl_2: 2M \, NH3 \, in \, MeOH = 20:1)$ and pure product was recrystalized from evaporated. The crude product was applied to short silica-gel column chromatography satNaHCO3 and brine. The separated organic layer was dried over NaSO4 and Et20/ CH2Cl3. White powder. 1.62g(69%). C/MS: m/z 237(M+1) 2

Synthesis of (2)

13

This compound was synthesized according to the method described in the preparation of

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Synthesis of (3)

500mg of compound (1) (2.12mmol) was dissolved in 5.0ml of CH₂Cl₂ and cooled to 0 °C. 10.0ml of BBr3 1.0M in CH₂Cl₂ (10mmol) was added slowly and stirred at 0°C for 11h. McOH was added to quench the reaction and 4.0ml of 5NaOHaq. was added. The mixture was stirred at 0°C for 10min. CH₂Cl₂ layer was separated. The water layer was acidified slowly PH=14→2 and extracted with CH₂Cl₂ for each step. The water layer was concentrated in vacuo, filtered off NaCl. The filtrate was made to PH=10 stepwise and extracted with CH₂Cl₂ each step. All of these extractions were combined together, dried over NaSO4 and evaporated to give the product 301mg (64%). LC/MS: mfz 223(M+1)

Synthesis of (4)

This compound was synthesized according to the method described in the preparation of (3).

Synthesis of (5)

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52mg of compound (3) (0.23mmol), 57mg of 3-diethylaminopropanol (0.28mmol) and 73mg of Triphenylphosphine (0.28mmol) were dissolved in 2.0ml of dry THF. The air was replaced to N₂ gas. 37mg of Diisopropyl-azodicarboxylate (0.28mmol) in 0.5ml of THF was added to this reaction mixture and stirred at room temparature for overnight.

20 The reaction mixture was concentrated and applied to SCX column, washed by McOH. The crude product was eluted with 2M NH3 in McOH. This crude product was applied to silica-gel column chromatography (CH₂Cl₂: 2M NH3 in McOH = 20:1) to give the product. 48mg (62%). LC/MS: m/z 336(M+1)

Synthesis of (6)

z

This compound was synthesized according to the method described in the preparation of

Synthesis of (7)

30 3.0ml of Litium aluminium hydride 1.0M in THF (3.0mmol) was placed in flask and the air was replaced to N2gas. 43mg of compound (5) (0.13mmol) in 2.0ml of THF was added slowly into the flask and stirred under reflux for 2h. The reaction mixture was

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allowed to cool to room temperature and water was added to quench the reaction. The organic layer was decanted. The water layer was extracted with CH₂Cl₂ (3 times) and all organic layers were combined together. This solution was dried over NaSO4 and evaporated. The crude product was applied to silica-gel column chromatography (CH₂Cl₂: 2M NH3 in MeOH = 20:1) to give the product. 19mg (46%). LCMS: m/z 322(M+1)

Synthesis of (8)

This compound was synthesized according to the method described in the preparation of (7).

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Synthesis of (10) 100mg of compound (9) (0.50mmol) and 116mg of (R)(-)-1-(2-pyrrolidinylmethyl)pyrrolidine (0.75mmol) were dissolved in 5.0ml of 5%AcOH in CH₂Cl₂ and 310mg of MP-cyanoborohydride (mmo/g =2.42, 0.75mmol) was added in the reaction vial. The vial was capped by Teflon cap and shaken at 60°C for overnight. The reaction mixture was filtered and the filtrate was concentrated under N2 gas. The crude product was applied to silica-gel column chromatography (CH₂Cl₂: 2M NH3 in McOH = 20:1) to give the product. 143mg (85%). LC/MS: m/z 337(M+1)

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Synthesis of Example 261

20 65 mg of compound (10) (0.19mmol) and 50mg of piperidine (0.58mmol) were put into
4.0ml vial and 2.0ml of THF and 10mg of NaI were added to the vial. The vial was
capped by Teflon cap and heated at 100°C for 3days. The reaction mixture was
concentrated under N2gas and applied to silica-gel column chromatography (CH₂Cl₂: 2M
NH3 in MeOH = 20:1) to give the product. 38mg (51%). LC/MS: m/z 386(M+1)

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Synthesis of (15)

mixture was diluted with CH2Cl2 and washed by brine. The crude product was applied to and stirred at 70°C for 1h under N2 gas. The excess acid chloride was removed in vacuo. (4.17mmol) were dissolved in 10ml of CH2Cl2 and cooled to 0°C. Acid chloride solution The residue was dissolved in 1.0ml of CH₂Cl₂ to make acid chloride solution. 643mg of 813mg of compound (14) (98536) (3.8mmol) was dissolved in 5.0ml of thionyl chloride was added to this mixture at 0°C and stirred at room temperature for 2h. The reaction (S)(+)-1(2-pyrrolidinylmethyl)pyrrolidine (4.17mmol) and 421mg of triethylamine silica-gel column chromatography (CH₂Cl₂: 2M NH3 in MeOH = 10.1) to give the product. 1.13g (85%) LC/MS: m/z 351(M+1)

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Synthesis of Example 209

15 This compound was synthesized according to the method described in the preparation of Example 261.

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Synthesis of (18)

column chromatography (CH₂Cl₂: 2M NH3 in McOH = 20:1) to give the product. 1.64g 60°C for 1h. Almost of MeOH was removed in vacua. The residue was dissolved in 1.17g of Na(51mmol) was dissolved in 200ml of McOH and 6.48g of methyl p-hydroxy (119.6mmol). The reaction mixture was stirred at room temperature for 2h and stirred at water and acidified by cHCl to PH=1.0 and extracted with CH₂Cl₂. The separated organic layer was dried over NaSO4 and evaporated. The crude product was applied to silica-gel (17%). NMR (DMSO); 7.84(d, 2H, 1=5.9Hz), 6.91(d, 2H, 1=5.9Hz), 4.02(t, 2H, benzoate(17) (42.5mmol) was added followed by 20.52g of 1-bromo 4-chlorobutane J=5.8Hz), 3.69(t, 2H, J=6.4Hz), 1.85(m, 4H) 2

NaOHaq, was added. The reaction mixture was stirred at room temperature for overnight. 1.14g of compound (19) (4.44mmol) was dissolved in 15ml of MeOH and 10ml of 5N Synthesis of (20)

The reaction mixture was evaporated. The residue was dissolved in water and acidified by cHCl to PH=1.0. This solution was extracted with CH₂Cl₂, dried over NaSO4 and evaporated. The pure product was recrystalized from Hexanel CH₂Cl₂. 829mg (77%) NMR (DMSO); 8.05(d, 2H, J=8.9Hz), 6.93(d, 2H, J=8.9Hz), 4.05(t, 2H, J=6.3Hz), 3.57(t, 2H, J=6.8Hz), 1.86(m, 4H), 1.65(m, 2H) 15

CHCl₃:CH₃CN:IBuOH (5:1:1) 1. —-DCC, HOBI

(101)

1-(4-(3-Piparidin-1-yl-propoxy)-phenyll-butan-1-one To a 20 mL. vial was placed keto-phenol (500 mg, 3 mmol), CsCO₃ (1.98 g, 6 mmol), KI (454 mg, 3 mmol) and chloropropylpiperdine (64 mg, 3.3 mmol). Dioxane added and the reaction was heated to 90 degrees overnight on a J-KEM heater/shaker block. The The material was purified by Biotage utilizing 4:1 EtOAc:McOH to afford (201) as a reaction was then quenched with water, extracted into DCM and dried over Na2SO4. orange oil (880 mg, 99%). Mass spec hit M+1, 290; LCMS >95% @ 230 nm and ELSD.

> To a 4 mL vial was placed 101 (28.5 mg, 0.1 mmol), resin-bound DCC (170 mg, 0.16 overnight. In the morning, PS-trisamine (134 mg, 0.4 mmol, 3.0 mmol/g) was added and the reaction again allowed to rotate overnight to scavenge excess carboxylic acid and HOBt. Filtration, washing with DCM/MeOH and concentration afforded a orange foam. Filtration through a short pipet column provided 25 mg (83%) of an yellow solid, 629304. mmol, 0.94 mmol/g), HOBt (16 mg, 0.12 mmol), amine (13 uL, 0.08 mmol) and a 5:1:1 mixture of CHCl3:CH3CN:tBuOH. The vial was agitated by means of a lab quake shaker Mass spec hit M+1, 386; LCMS >95% @ 230 nm and ELSD. A substantially analogous procedure was employed for the array synthesis of Examples:

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Observed Mass	361	361	389	401	386	386	401	372	400	360	340	346	360	360	386	350
Example #	41	42	44	43	130	131	132	133	144	150	151	152	153	154	155	172

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To a 20 mL vial was placed (102) (300 mg, 1 mmol), diamine (120 mg, 1.14 mmol), MP-CNBH₃ (2.4 g, 6.22 mmol) and a 9:1 CHCl₃:HOAc solution. The reaction was heated to 50 degrees overnight on a J-KEM heater/shaker block. The reaction was filtered, washed with DCM/MeOH. The material was then subjected to preparative HPLC purification to afford 29 mg (3%) of analytically pure example 94. as a white solid. Mass spec hit M+1, 362; LCMS >98% @ 230 nm and ELSD. Example 192 can be made by a substantially analogous procedure, Observed mass 360. The following examples are made by a substantially analogous procedure:

(M+1)	320	246 M-87	38	35	336	272 M-67	258 M-87	8	ğ	88	88
Example	7	8		81	88	68	8	<u> </u>	23	· \$	8
Product Name	N-[8 (3-Dimethylamino-propoxy)-1,2,3,4-tetrahydro- naphthalen-1-yl]-N,N-dimethyl-ethane-1,2-diamine	N-(6-(3-Dimethylamino-2-methyl-propoxy)- 1,2,3,4-tetrahydro-naphthalen-1-yll- N,N-dimethyl-ethane-1,2-diamine	N,A-Dimethy-M-(6-(1-methy-plperkin-3- ymethoxy)-1-2,3,4-tetuahydro-naphbalon- 1-yl-ethane-1,2-diamine	N-{1-14-(3-Dimethylamino-2-methyl-propoxy)- phenyl-propyl-v , N-dimethyl- ethane-1,2-diamine	N-{1-{4-{3-Dimetrylamino-2-metryl-propoxy}- phenyl-butyl-N./N-dimetryl- ethane-1,2-diamine	N.N-Dmethyl-N-(6-(3-pbendin-1-yl-propoxy)- 1,2,3,4-tetrahydro-naphthalen-1-yll-ethane- 1,2-diamine	N.N.Dimetry: N-{6-{2-piperidin-1-y-ethoxy}- 1,2,3,4-lefrahydro-naphthalen-1-y }-ethane- 1,2-diamine	N, N-Dinebtyl-N-(1-(4-(3-piperdin-1-yf-propoxy)- phemyl-propyl-ethane-1,2-dlamine	N.N-Dimethyl-N-(1-(4-(2-pipendin-1-ye-ethoxy)- phenyl-butyl)-ethane-1,2-diamine	N-{1-{4-(3-Dimethylamino-propoxy)-pheny(l-buty)- N,N-dimethyl-ethane-1,2-diamine	N,M-Dimethyl-N-(1-(4-(2-pipendin-1-yl-ethoxy)-phenyl)-butyl)-ethane-1,2-dismine
Phemy Ketone											* E

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Examples 135, 14, 126 6

NaBH4, MeOH

To a 10 mL round-bottom flask was added (102) (280 mg, 0.96 mmol) and dry MeOH (5 mL). Then, NaBH₄ (74 mg, 1.93 mmol) was added at room temperature. After 1 hour, the reaction was then quenched with water, extracted into DCM and dried over Na₂SO₄. The material was purified by Biotage utilizing 4:1 EtOAc:MeOH to provide 270 mg (98%) of a white solid. Mass spec hit M+1, 292; LCMS >98% @ 230 nm and ELSD. Examples 14 and 126 are made by a substantially analogous procedure. Observed mass: Example 14 = 321, Example 126 = 375.

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Example 142

Example 142

To a round-bottom flask, equipped with stir bar and septum, was placed (103) (300 mg, 1.03 mmol), KJ (230 mg, 1.54 mmol) and NaH (78 mg, 95%dry, 3.09 mmol). Then, dry DMF (20 mL, 0.5 M) was added via syringe followed by chloroethylpiperidine (285 mg, 1.54 mmol). The reaction was allowed to stir at 50 degrees overnight. In the moming, the reaction was quenched with water, extracted into EtOAc (3 x 20 mL) and dried over brine. Column chromatography in 9:1 DCM:MeOH afforded 631934 an yellow oil (300 mg, 79%). Mass sec hit M+1, 404; LCMS >95% @ 230 nm and ELSD.

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Example 246
NaH (95%)
DMF
DMF
0-RT

3-Piperidinylpropanol (3.56g, 25 mmoles) in 4 ml DMF was added to a slurry of sodium hydride in 10 ml DMF at 0 C., and the reaction was stirred at 0 C. for 0.5 hr. The 4 fluorobenzonitrile in 6 ml was added at 0 C. The reaction was stirred at 0 C for 1 hr. and at RT overnight. Water and ether were carefully added. Separated the ether layer and

at RT overnight. Water and ether were carefully added. Separated the ether layer and extracted with water five times. The ether extract was dried over sodium sulfate, filtered and evaporated to give 6.0g(0.0246 mmoles, 98.4% yield). LCMS 1.61 min @254.0 nm 95.2%; @220.0 nm 89.5%; ELSD 1.71 min 100%; MS 1.59 min M + 1 = 245 good for product (104).

Example 246

The nitrile(6.0g, 0.0246 mmoles) in 250 ml 2B EtOH with 2.5 g RaNi was hydrogenated at 80 C. for 8 hrs. Filtration and evaporation yielded 5.4 g oil(88.4 yield).

Example 217

The 1-hydroxybenzotriazole hydrate(13.5 mg, 0.1 munole),1-piperidinepropionic acid(18.1 mg, 0.115 mnole), amine(248 mg, 0.1 mmole), polystyrene-carbodiimide(125 mg, 0.15 mmoles) and 2.5 ml chloroform, acetonitrile, t-butanol(5:1:1) in a 4 ml vial were rotated for four days. Polystyrene-trisamine(93.7 mg, 0.4 mmoles) was added and the reaction was rotated overnight. Filtered reaction through filter cartridge and evaporated to give 37.5 mg, 0.0967 mmole, 96.7% yield. LCMS ELSD 1.42 min 100%, MS 1.21 min M + 1 = 388 good for product.

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Example 1

The solution of diisopropylazodicarboxylate(3.93 ml, 20 mmoles) in 20 ml anhydrous THF was added dropwise with stirring to the cold solution of 4-

hydroxyacetophenone(2.18 g. 16 mmoles), 3-diethylaminopropanol(2.23 ml, 15 mmoles) and triphenylphosphine(4.98 g., 19 mmoles) in 50 ml anhydrous THF over 45 minutes. The reaction was stirred in an ice bath for one hour and at room temperature for 18 hours. The solvent was evaporated and ether was added. This solution was extracted with dilute HCI(1.0 N) four times. These combined acidic extracts were extracted with ether, basified with a NaOH solution and extracted with ether times. These combined

oethereal extracts were dried over sodium sulfate, filtered and evaporated to give 3.41 g oil. LCMS 1.53 min @254.0 nm 97.4%; ELSD 1.59 min 91.1%; MS 1.58 min M+1=250 good for product (105).

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In a 7 ml vial with cap, 4-(3-diethylaminopropyloxy)acetophenone(0.47 g, 0.19 mmoles),
N-(2-aminoethyl)morpholine(0.039 ml, 0.3 mmoles) and macroporus
cyanoborohydride(169 mg, 0.4 mmoles) in 2 ml dichloromethane with 0.2 ml glacial

acetic were heated on shaker at 550 for 18 hours. Purified with a 3 ml extrelut cartridge

20 hydrated with 3 ml water. The reaction solution was added and the cartridge was rinsed with dichloromethane(5 ml). The product was eluted with 10% triethylamine/dichloromethane. LCMS 1.14 min @254.0 nm 95.6%; @230.0 nm 95.3%; I.20 min ELSD 95.3%; MS 1.14 min M+1=364 good for product.

with 3 ml water. The reaction solution was added and the cartridge was rinsed with In a 7 ml vial with cap, 4-(3-diethylaminopropyloxy)benzaldehyde(0.59 g, 0.25 mmoles), cyanoborohydride(210 mg, 0.5 mmoles) in 3 ml dichloromethane with 0.3 ml glacial acetic were heated on shaker at 40° briefly. Purified with 3 ml extrelut cartridge hydrated dicloromethane(5 ml). The product was eluted with 10% triethylamine/dichloromethane. LCMS 1.14 min ELSD 95.3%; MS 1.09 min M+1=350 good for product Example 62. mmoles) and 0.375 N-(2-aminoethyl)morpholine(0.049 · ml,

364 348 308 336 337 391 336

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363 362 359 336 376	(901)
231 24 25 26 27	Example 62 TPP DIAD OPEN OFFT OFFT

hydroxybenzaldehyde(1.95 g, 16 mmoles), 3-diethylaminopropanol(2.23 ml, 15 mmoles) The solution of diisopropylazodicarboxylate (3.93 ml, 20 mmoles) in 20 ml anhydrous THF was added dropwise with stirring to the cold solution of 4-

and triphenylphosphine (4.98 g, 19 mmoles) in 50 ml anhydrous THF over 45 minutes.

The solvent was evaporated and ether was added. This solution was extracted with dilute oil. LCMS 1.47 min @254.0 nm 97.0%; ELSD 1.53 min 95.4%; MS 1.48 min M+1=236 The reaction was stirred in an ice bath for one hour and at room temperature for 18 hours. ethereal extracts were dried over sodium sulfate, filtered and evaporated to give 3.71 g basified with a NaOH solution and extracted with ether three times. These combined HCl(1.0 N) four times. These combined acidic extracts were extracted with ether, good for product. 13

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Observed Mass	320	334	294	348	348	322	363	377	322	349	348	345	322	362	¥.	376	348	320	420	410	334	334
Example	629	63	47	84	49	20	51	22	61	53	¥	2	17	72	73	29	74	\$	113	114	101	103

Example 45
Cs₂CO₃

Kl
No Dioxane, water
BS9

C107)

4-Hydroxybenzaldehyde(2.44g, 20 mmoles), N-(3-Chloropropyl)pipendine hydrochloride, cesium carbonate(19.7 g, 60 mmoles) and potassium iodide in 14 ml dioxane with 0.7 ml water were stirred at 85º for 8 hours and at room temperature for 16 hours. Evaporated the decanted supernatant, added water to both (evaporated supernatant and solid) and extracted three times with ether. These combined ethereal extracts were washed three times with water, dried over sodium sulfate, filtered and evaporated to give 7.8 g oil. LCMS 1.48 min @254.0 nm 99.4%; @230.0 nm 89.6%; 1.51 min ELSD 99.4%; MS 1.49 min M+1=248 good for product. 300 mHz NMR(CDCl3) good for structure (107).

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In a 7 ml vial with cap, 4-[(3-N-piperidinyl)propyloxylbenzaldehyde(0.062 g, 0.25 mmoles), N-(2-aminoethyl)morpholine(0.049 ml, 0.375 mmoles) and macroporus cyanoborohydride(210 mg, 0.5 mmoles) in 3 ml dichloromethane with 0.3 ml glacial acetic were heated on shaker at 40°. The reaction was shaken at room temperature for 16 hours and at 40° for one hour. Purified with 3 ml extrelut cartridge hydrated with 3 ml water. The reaction solution was added and the cartridge was finsed with dicloromethane(5 ml). The product was eluted with 10% triethylamine/dichloromethane. LCMS 1.13 min @230.0 nm 97.3%; 1.19 min ELSD 98.5%; MS 1.13 min M+1=362 good for product Example 45.

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Example 100

Dimethyl-(3-(4-[1-(2-piperidin-1-yl-ethylamino)-ethyl]-phenoxyl-propyl)-amine
To a 20 mL vial was placed (108) (42 mg, 0.19 mmol), amine (37 mg, 0.29 mmol), MPCNBH₃ (190 mg, 0.45 mmol, 2.37 mmol/g) and a 9:1 CHCl₃:HOAc solution. The
reaction was heated to 50 degrees overnight on a J-KEM heater/shaker block. The
reaction was filtered, washed with DCM/MeOH. The material was then subjected to
preparative HPLC purification to afford 5.8 mg (9%) example 100. As a clear oil. Mass
spec hit M+1, 334; LCMS >89% @ 214 nm.

In a procedure substantially similar to that for synthesis if Example 100, the following examples are made:

Activizations Amino Product Name

| Main | Dimension | 13 | Mis | Dimension | 13 | Mis | Dimension | 13 | Dimension | 14 | Dimension | 15 | Di

N-{1-[4-(3-Diethylamino-propoxy)-phenyl]-ethyl}-N-(2-dimethylamino-ethyl)-C-phenyl-methanesulfonamide. To a 4 ml vial was placed N-{1-[4-(3-Diethylamino-propoxy)-phenyl]-ethyl}-N'N-dimethyl-ethane-1,2-diamine (22 mg, 0.07 mmol), phenyl-methanesulfonyl chloride (27 mg, 0.14 mmol), PS-DMAP (93 mg, 1.48 mmol)g, and CH₂Cl₂ (1.5 ml). The vial was agitated by means of a lab quake shaker for 4 h. To the solution was added PS-Trisamine (100 mg, 3.3 mmol, 3.0 mmol/g) and the reaction was allowed to agitate overnight to scavenge excess methansulfonyl chloride. Filtration, washing with CH₂Cl₂ and concentrating afforded N-{1-[4-(3-Diethylamino-propoxy)-phenyl]-cHyl]-N-(2-dimethylamino-ethyl)-C-phenyl-methanesulfonamide. Mass spec hit M+1, 476: LCMS >93% @ 230 nm and ELSD.

MS (M+1)	462	468	468
Example	8	8	31
Product Name Es	N-{1-{4-(3-Diethylamino-propoxy}-phenyl}-ethyl}- N-{2-dimethylamino-ethyl}-benzenesutionamide	Thiophene-2-suffonic acid (1-[4-(3-diethylamino-propoxy)-phenyl-ethyl-(2-dimethylamino-ethyl)-amide	2.2.2-Tiffuoro-ethanesulionic acid (1-44/3-diethyfamino-propoxy)-phenylj-ethyl- (2-dimethyfamino-ethyl)-amide
Sulfonyl Chloride	lo-so-ci	12 ² OS-{}	ات، دەمىرا

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compounds of Formula I and Formula II were prepared. Structural figures for representative examples of Formula I and Formula II are shown the following pages. Utilizing the procedures provided herein, in addition to methods known in the $\operatorname{\mathsf{art}}$,

Observed Mass	336	321.2	
Structure	5 2 5	£	
Example Number		. 7	m

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386	375.3	275.2	371.4	415.2	385.2
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	290	291	292	293	294

375.3	275.3	371.4	303.3	415.3	385.3	371.4
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307	308	309	310	311	312	313

374	415.3	418.4	433.2	433.2	303.3
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320	331	322	323	25

389.3	317.2	389.3	385.3	428	
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The compound of Formula I is preferably formulated in a unit dosage form prior to administration. Therefore, yet another embodiment of the present invention is a

pharmaceutically acceptable carriers, diluents or excipients.

pharmaceutical composition comprising a compound of Formula I and one or more

The present pharmaceutical compositions are prepared by known procedures using well-known and readily available ingredients. In making the formulations of the present invention, the active ingredient (Formula I compound) will usually be mixed with a carrier, or diluted by a carrier, or enclosed within a carrier which may be in the form of a capsule, sachet, paper or other container. When the carrier serves as a diluent, it may be a solid, semisolid or liquid material that acts as a vehicle, excipient, or medium for the active ingredient. Thus, the compositions can be in the form of tablets, pills, powders, lozenges, sachets, cachets, elixirs, suspensions, emulsions, solutions, syrups, aerosol (as a solid or in a liquid medium), soft and hard gelatin capsules, suppositories, sterile injectable solutions and sterile packaged powders.

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Some examples of suitable carriers, excipients, and diluents include lactose, dextrose, sucrose, sorbitol, mannitol, starches, gum acacia, calcium phosphate, alginates, tragacanth, gelatin, calcium silicate, microcrystalline cellulose, polyvinylpytrolidone, cellulose, water syrup, methyl cellulose, methyl and propylhydroxybenzoates, talc, magnesium stearate and mineral oil. The formulations can additionally include lubricating agents, wetting agents, emulsifying and suspending agents, preserving agents, sweetening agents or flavoring agents. The compositions of the invention may be formulated so as to provide quick, sustained or delayed release of the active ingredient after administration to the patient.

The compositions of the present invention may be formulated in sustained release form to provide the rate controlled release of any one or more of the components or active ingredients to optimize the therapeutic effects, i.e., antihistaminic activity and the like.

Suitable dosage forms for sustained release include layered tablets containing layers of varying disintegration rates or controlled release polymeric matrices impregnated with the active components and shaped in tablet form or capsules containing such impregnated or encapsulated porous polymeric matrices.

Liquid form preparations include solutions, suspensions and emulsions. As an example may be mentioned water or water-propylene glycol solutions for parenteral injections or addition of sweeteners and opacifiers for oral solutions, suspensions and emulsions. Liquid form preparations may also include solutions for intranasal administration.

Aerosol preparations suitable for inhalation may include solutions and solids in powder form, which may be in combination with a pharmaceutically acceptable carrier such as inert compressed gas, e.g. nitrogen.

For preparing suppositories, a low melting wax such as a mixture of fatty acid glycerides such as cocoa butter is first melted, and the active ingredient is dispersed homogeneously therein by stirring or similar mixing. The molten homogeneous mixture is then poured into convenient sized molds, allowed to cool and thereby solidify.

Also included are solid form preparations which are intended to be converted, shortly before use, to liquid form preparations for either oral or parenteral administration Such liquid forms include solutions, suspensions and emulsions.

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The compounds of the invention may also be deliverable transdermally. The transdermal compositions may take the form of creams, lotions, aerosols and/or emulsions and can be included in a transdermal patch of the matrix or reservoir type as a re conventional in the art for this purpose.

20 Preferably the compound is administered orally.

Preferably, the pharmaceutical preparation is in a unit dosage form. In such form, the preparation is subdivided into suitably sized unit doses containing appropriate quantities of the active components, e.g., an effective amount to achieve the desired purpose.

The quantity of the inventive active composition in a unit dose of preparation may be generally varied or adjusted from about 0.01 milligrams to about 1,000 milligrams, preferably from about 0.01 to about 950 milligrams, more preferably from about 0.01 to about 500 milligrams, and typically from about 1 to about 250 milligrams, according to the particular application. The actual dosage employed may be varied depending upon the patient's age, sex, weight and severity of the condition being treated. Such techniques

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are well known to those skilled in the art. Generally, the human oral dosage form containing the active ingredients can be administered 1 or 2 times per day.

More particularly, these compounds are selective histamine H3 receptor antagonists that have little or no affinity for histamine receptor GPRv53(H4R). As selective antagonists, conditions responsive to the inactivation of the histamine H3 receptor, including but not Compounds of Formula I are effective as histamine H3 receptor antagonists. the compounds of Formula I are useful in the treatment of diseases, disorders, or

limited to obesity and other eating-related disorders. It is postulated that selective antagonists of H3R will raise brain histamine levels and possibly that of other

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consequences. Although a number of H3R antagonists are known in the art, none have monoamines resulting in inhibition of food consumption while minimizing peripheral

proven to be satisfactory obesity drugs. There is increasing evidence that histamine plays an important role in energy homeostasis. Histamine, acting as a neurotransmitter in the many cell types and it binds to a family of G protein-coupled receptors (GPCRs). This hypothalamus, suppressed appetite. Histamine is an almost ubiquitous amine found in amily provides a mechanism by which histamine can elicit distinct cellular responses

density of expression of H3R was found in feeding center of the brain. A novel histamine peripheral white blood cells; only low levels have been identified in the brain by some investigators while others cannot detect it in the brain. However, any drug discovery primarily expressed in the brain, notably in the thalamus and caudate nucleus. High receptor GPRv53 has been recently identified. GPRv53 is found in high levels in effort initiated around H3R must consider GPRv53 as well as the other subtypes. 2

based on receptor distribution. Both the H1R and H2R are widely distributed. H3R is

transfected with cDNA coding for H3R to prepare membranes used for the binding assay. inhibition Scintillation Proximity Assay (SPA) based on a H3R binding assay using [3H] α methylhistamine as ligand. Stable cell lines, including but not limited to HEK can be The technique is illustrated below (Example 3) for the histamine receptor subtypes. The inventive compounds can readily be evaluated by using a competitive ĸ

Compounds of the invention of Formula I were tested for their ability to inhibit binding in Membranes isolated as described in Example 3 were used in a [35S]GTPxS functional assay. Binding of [35S]GTPXS to membranes indicates agonist activity,

protein was used per well in the SPA receptor-binding assay

Compounds of Formula I were tested for their ability to permit forskolin -stimulated the presence of agonists. Alternately, the same transfected cell lines were used for a cAMP assay wherein H3R agonists inhibited forskolin-activated synthesis of cAMP. cAMP synthesis in the presence of agonist.

Preparation of Histamine Receptor Subtype Membranes

A. Preparation H1R membranes

cDNA for the human histamine I receptor (H1R) was cloned into a mammalian Diagnostics Corporation). Transfected cells were selected using G418 (500 µ/ml). expression vector containing the CMV promoter (pcDNA3.1(+), Invitogen) and transfected into HEK293 cells using the FuGENE Transection Reagent (Roche

binding, cells were assayed in a SPA reaction containing 50mM Tris-HCL (assay buffer), confluent monolayers in 96-well dishes (Costar Clear Bottom Plates, #3632) by seeding wells with 25,000 cells and growing for 48 hours (37°C, 5% CO₂). Growth media was grown in 96-well dishes using a scintillation proximity assay (SPA) based radioligand Colonies that survived selection were grown and tested for histamine binding to cells binding assay. Briefly, cells, representing individual selected clones, were grown as removed and wells were rinsed two times with PBS (minus Ca2+ or Mg2+). For total pH 7.6, 1mg wheat germ agglutinin SPA beads (Amersham Pharmacia Biotech, 2 15

#RPNQ0001), and 0.8nM 3H-pyrilamine (Net-594, NEN) (total volume per well = 200µl). representing ~10 grams, were resuspended in 30ml assay buffer, mixed by vortexing, and specific binding. Plates were covered with FasCal and incubated at room temperature for 120 minutes. Following incubation, plates were centrifuged at 1,000rpm (~800g) for 10 centrifugation was repeated 2 more times. The final cell pellet was reusupened in 30ml scintillation counter. Several clones were selected as positive for binding, and a single Astemizole (10µM, Sigma #A6424) was added to appropriate wells to determine nonminutes at room temperature. Plates were counted in a Wallac Trilux 1450 Microbeta centrifuged (40,000g at 4°C) for 10 minutes. The pellet resuspension, vortexing, and and homogenized with a Polytron Tissue Homogenizer. Protein determinations were done using the Coomassie Plus Protein Assay Reagent (Pierce). Five micrograms of clone (H1R40) was used to prepare membranes for binding studies. Cell pellets, 2 23 8

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B. Preparation H2R membranes

cDNA for the human histamine 2 receptor was cloned, expressed and transfected into HEK 293 cells as described above. Histamine binding to cells was assayed by SPA (Amersham Pharmacia Biotech, #RPNQ0001), and 6.2nM3H-tiotidine (Net-688, NEN) described above. For total binding, cells were assayed in a SPA reaction containing (total volume per well = 200µl). Cimetidine (10µM, Sigma #C4522) was added to 50mM Tris-HCl (assay buffer), pH 7.6, 1mg wheat germ agglutinin SPA beads appropriate wells to determine non-specific binding.

Several clones were selected as positive for binding, and a single clone (H2R10) was used to prepare membranes for binding studies. Five micrograms of protein was used per well in the SPA receptor-binding assay.

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C. Preparation of H3R membranes

in Example 1, above. Transfected cells were selected using G418 (500 µ/ml), grown, and prepare membranes for binding studies described above. Five micrograms of protein was cDNA for the human histamine 3 receptor was cloned and expressed as described #RPNQ0001), and 1nM (3H)-n-alpha-methylhistamine (NEN, NET1027) (total volume per well = 200µl). Thioperimide was added to determine non-specific binding. Several tested for histamine binding by the SPA described above. For total binding, cells were assayed in a SPA reaction described above containing 50mM Tris-HCL (assay buffer), clones were selected as positive for binding, and a single clone (H3R8) was used to pH 7.6, 1mg wheat germ agglutinin SPA beads (Amersham Pharmacia Biotech, used per well in the SPA receptor-binding assay. 15

the H3 receptor greater than 200 n.M. Most preferred compounds of the invention exhibit receptor greater than 1 u.M. Preferred compounds of the invention exhibited affinity for All compounds set forth in examples 1 to 322 exhibited affinity for the H3 affinity for the H3 receptor greater than 20 nM.

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D. Preparation of GPRv53 Membranes

cDNA for the human GPRv53 receptor was cloned and expressed as described in selected. HEK293 GPRv53 50 cells were grown to confluency in DMEM/F12 (Gibco) Example 1, above. Transfected cells were selected, tested for histamine binding, and 8

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96 well dishes with 3 nM (3H) Histamine and compounds in binding buffer for 2 hours at tissuemizer in binding buffer, 50 mM Tris pH 7.5. Cell lysates, 50 ug, were incubated in room temperature. Lysates were filtered through glass fiber filters (Perkin Elmer) with a supplemented with 5 % FBS and 500 ug/ml G418 and washed with Delbecco's PBS (Gibco) and harvested by scraping. Whole cells were homogenized with a Polytron Tomtec cell harverster. Filters were counted with melt-on scintillator sheets (Perkin Elmer) in a Wallac Trilux 1450 Microbeta Scintillation counter for 5 minutes.

Pharmacological Results

CAMP ELISA 2

temperature. Then 50 µl of cell culture medium containing 20 µM Forskolin (Sigma) was (Sigma) and incubated for 20 minutes at room temperature. Antagonist were added in 50 replaced with 50 µl cell culture medium containing 4 mM 3-isobutyl-1-methylxanthine 50,000 cells/well and grown overnight in DMEM/F12 (Gibco) supplemented with 5 % medium was removed and cells were lysed in 0.1M HCl and cAMP was measured by added to each well and incubated for 20 minutes at room temperature. Tissue culture HEK293 H3R8 cells prepared as described above were seeded at a density of added to the wells in 50 µl cell culture medium and incubated for 5 minutes at room R (-)α methylhistamine (RBI) at a dose response from 1x10⁻¹⁰ to 1x10⁻³ M was then μl cell culture medium and incubated for 20 minutes at room temperature. Agonist FBS and 500 ug/ml G418. The next day tissue culture medium was removed and ELISA (Assay Designs, Inc.). 20 34 2

[35S] GTP y [S] Binding Assay

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Antagonist activity of selected compounds was tested for inhibition of [35S] GTP temperature in 20 mM HEPES, 100 mM NaCl ,5 mM MgCl₂ and 10 uM GDP at pH 7.4 expressing HEK293 cell line (20 ug/well) and GDP were added to each well in a volume of 50 µl assay buffer. Antagonist was then added to the wells in a volume of 50 µl assay y [S] binding to H3R membranes in the presence of agonists. Assays were run at room in a final volume of 200 ul in 96-well Costar plates. Membranes isolated from H3R8buffer and incubated for 15 minutes at room temperature. Agonist R(-)alpha

and incubated for 5 minutes at room temperature. GTP y [35S] was added to each well in addition of 50 µl of 20 mg/ml WGA coated SPA beads (Amersham). Plates were counted inhibited more than 50% of the specific binding of radioactive ligand to the receptor werr concentration of 100 nM were then added to the wells in a volume of 50 µl assay buffer in Wallac Trilux 1450 Microbeta scintillation counter for 1 minute. Compounds that methylhistamine (RBI) at either a dose response from $1x10^{-10}$ to $1x10^{-5}$ M or fixed a volume of 50 µl assay buffer at a final concentration of 200 pM, followed by the serially diluted to determine a K[i](nM). The results are given below the indicated compound.

Table 1

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Example 2

4. Example 1

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antagonists also bound H4R. As demonstrated in Table 2, example 131 and example 250 To investigate the selectivity of the antagonists for the histamine receptors, a competitive did not inhibit binding H4R compare to H3R. To our knowledge, the study in Table 2 is determined. Importantly, the identification of H3R-specific antagonists that do bind the newly identified H4R was demonstrated. Until the present invention, most known H3R (structures given above) to selectively inhibit binding to H3R, H1R, H2 and H4R was binding assay described above was performed. The ability of example 131 and 250 the first demonstration of a H3R specific antagonist.

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Table 2 Ki (nM)

		,		
Compound.	H3R	H4R	HIR	H2
Example 131 1.05	1.05	≥ 20,000	≥ 20,000	≥ 20,000
Example 250 0.37	0.37	≥ 20,000	1022	6011

unexpectedly improved pharmacokinetic properties. Male Sprague Dawley Rats (n=3 per literature generally have very poor pharmacokinetic properties (see J. Apelt, et al, J. Med. dose arm) were separately dosed with 3 mg/kg iv or 10 mg/kg po of compound examples. 131 and 271 (vehicle: 5% ethanol/water or water respectively; dose volume: 1 mL/kg iv, 10 mL/kg po). Approximately 0.5 mL of blood was collected in hepann collection tubes Non-imidazole containing histamine H3 receptor antagonists disclosed in the Chem. 2002, 45, 1128-1141). Compounds of this invention have markedly and at multiple time points over an 8 or 24-hour period for examples 131 and 271 2

respectively, and the samples were analyzed using LC/MS/MS. In this manner compound have an oral bioavailability of 69% (AUC 0-24hr; posiv ratio) and an oral half-life of 71.9 example 131 was found to have an oral bioavailability of 58% (AUC 0-24hr; posiv ratio) and an oral half-life of 10.4 ± 4.2 hours (±SEM). Compound example 271 was found to ± 3.3 hours (±SEM). 15

characteristics of the present invention, and without departing from the spirit and scope various usages and conditions. Thus, other embodiments are also within the claims. thereof, can make various changes and modifications of the invention to adapt it to From the above description, one skilled in the art can ascertain the essential

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WHAT IS CLAIMED IS:

A compound structurally represented by Formula I

or pharmaceutically acceptable salts thereof wherein:

X is O, NR⁷ or S;

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R1 is hydrogen,

 $C_1\text{-}C_8$ alkyl optionally substituted with 1 to 4 halogens,

(CHR⁵)_n-C₃-C₇ cycloalkyl,

(CHR⁵)_n aryl,

(CHR⁵)_n heteroaryl, or

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(CHR5)n-0(CHR5)n-aryl;

 \mathbb{R}^2 is independently \mathbb{R}^1 , or

 $\mathsf{COR}^1 \cdot \mathsf{or}\ \mathsf{cyclized}\ \mathsf{with}\ \mathsf{the}\ \mathsf{attached}\ \mathsf{nitrogen}\ \mathsf{atom}\ \mathsf{at}\ \mathsf{the}\ R^1\ \mathsf{position}\ \mathsf{to}\ \mathsf{form}\ \mathsf{a}\ \mathsf{4},$ 5, or 6 member carbon ring, wherein one of said carbons is optionally replaced by one of O, S, NR 1 or CO, or wherein the ring formed by R 1 and R 2 is optionally substituted one to two times with C1-C4 alkyl; 2

 R^3 is independently $C_5\text{--}C_7$ cycloalkylene, or $C_l\text{--}C_4$ alkylene optionally substituted;

R4 is hydrogen,

halogen,

C₁-C₄ alkyl,

(CHR5)_n-C₃-C₇ cycloalkyl,

(CHR⁵)_h aryl,

(CHR⁵)_n heteroaryl,

(CHR⁵)_n-O(CHR⁵)_n-aryl or

COo

cyclized with R5 to from a cyclopropyl ring;

2

R⁵ is hydrogen, or

C₁-C₄ alkyl;

15 R⁶ is hydrogen,

halo or

cyclized with the attached carbon atom at the R⁵ position to form a 5 to 6 member

carbon ring,

cyclized with the attached carbon atom at the R⁷ position to form a 5 to 6 member

heterocyclic nng or

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R7 is hydrogen,

C1-C8 alkyl optionally substituted with 1 to 4 halogens,

(CHR5)n-C3-C7 cycloalkyl,

(CHR⁵)_n aryl,

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(CHR⁵)_n-O(CHR⁵)_n-aryl, (CHR5)_n heteroaryl,

SO2R1 or

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Cyclized with attached carbon on \mathbb{R}^8 to from a 5, 6, or 7 membered carbon ring	ia 5, 6, or 7 membered carbon ring			-CONR ¹ R ²		
optionally substituted with R9, CF3, or CN, optionally one of the said carbons is replaced	lly one of the said carbons is replaced			-NHSO ₂ R ¹ ,		
by N, NR ¹ , CO;				-NO ₂ ,		
				-co ₂ R ¹ ,		•
K ^o is hydrogen, s band			, s	$-SO_2N(\mathbb{R}^1)_2$,		
C ₁ -C ₈ alkyl				-S(O) _n R ¹ ,		
-SO ₂ R ⁹ .				-OCF ₃		
018 00				-CH2SR5,		
6400			R ¹⁰ i	R ¹⁰ is hydrogen,	•	
-CO RY,			01	halogen,		
-CONH RIG:				C1-C8 alkyl optionally substituted with 1 to 4 halogens,	ituted with 1 to 4 halogens,	
0.0		٠		C3-C7 cycloalkyl,		
r is nydrogen,				aryl,		
inarogen, Ci-Co alkel entionally substituted with 1 to 4 halosens	4 halogene			CH ₂ aryl,		
Clare cortoalty	i i ai Ogens,		15	heteroaryl,		
(3 C) cyclomings,		÷.		heterocycle,		
aryl,				-cor1,		
CH2 aryl,				CONR ¹ R ² ,		
heterocycle,				-so ₂ r ¹ ,		
-O(CHR ⁵) _n -aryl,			20	-N(R ¹) ₂ ,		
-cor1,				-NR1 R2,		
-conr ¹ R ² ,				-CH2NR1 R2,		
-SO ₂ R ¹ ,				-CONR1 R2		
-OR1,	·			-co ₂ R ¹ ,		
-N(R ¹) ₂ ,			25	-SO ₂ N(R ¹) ₂ ,		
-NR ¹ R ² ,				-S(O) _n R ¹ ,		
-CH ₂ NR ¹ R ² ,				-CH2SR ⁵ ,		

A compound of claim 1, structurally represented by Formula II

or pharmaceutically acceptable salts thereof where:

X is O, N or S;

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R^{1'} is hydrogen,

C1-C8 alkyl (optionally substituted with 1 to 4 halogens or C1-C4 alkyls),

(CHR5')n-C3-C7 cycloalkyl,

(CHR^{5'})_n aryl,

(CHR^{5'})_n heteroaryl, or

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(CHR^{5'})_n-O(CHR^{5'})_n-aryl;

 \mathbb{R}^{2} is independently \mathbb{R}^{1} , or

member carbon ring (optionally one of said carbons is replaced by one of O, S cyclized with the attached nitrogen atom at the R1' position to form a 5 to 6

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R3' is independently C1- C4 alkyl;

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R4' is hydrogen,

halogen,

Cı-C4 alkyl,

(CHR5')n-C3-C7 cycloalkyl,

(CHR⁵)_n aryl,

(CHR^{5'})_h heteroaryl,

(CHR⁵)_n-O(CHR⁵)_n-aryl or

carbonyl;

10 R5 is hydrogen or C1-C4 alkyl;

R6' is hydrogen, or

cyclized with the attached carbon atom at the R5' position to form a 5 to 6 member carbon ring, or cyclized with the attached carbon atom at the \mathbb{R}^7 position to form a 5 to 6 member heterocyclic ring; 15

 \mathbb{R}^{7} is hydrogen,

(CHR5')_n-C₃-C₇ cycloalkyl,

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C1-C8 alkyl (optionally substituted with 1 to 4 halogens or C1-C4 alkyls),

(CHR^{5'})_n aryl,

(CHR^{5'})_n heteroaryl,

(CHR⁵′)_n-O(CHR^{5′})_{n-aryl}

25 R8' is hydrogen,

C1-C8 alkyl (optionally substituted with 1 to 4 halogens or C1-C4 alkyls),

C3-C7 cycloalkyl,

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$$-SO_2R^{1}$$
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or a pharmaceutically acceptable salt or solvate thereof.

8. A compound of claim 1 wherein the compound has the structure:

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or a pharmaceutically acceptable salt or solvate thereof.

9. A compound of claim 1 wherein the compound has the structure:

or a pharmaceutically acceptable salt or solvate thereof.

10. A compound of claim 1 wherein the compound has the structure:

or a pharmaceutically acceptable salt or solvate thereof.

11. A compound of claim 1 wherein the compound has the structure:

or a pharmaceutically acceptable salt or solvate thereof.

12. A compound of claim 1 wherein the compound has the structure:

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or a pharmaceutically acceptable salt or solvate thereof.

A compound of claim 1 wherein the compound has the structure:

or a pharmaceutically acceptable salt or solvate thereof.

 A pharmaceutical composition which comprises a compound of any of claims 1-14 and a pharmaceutically acceptable carrier. 15. A method of selectively increasing histamine levels in cells by contacting the cells with an antagonist of the histamine H3 receptor, said antagonists being a

compound of any of claims 1-14.

16. A method of selectively increasing histamine levels in cells by contacting the cells with an antagonist of the histamine H3 receptor, said antagonists being a

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17. A method of selectively increasing histamine levels in cells by contacting the cells with an antagonist of the histamine H3 receptor, said antagonists being a compound of Claim 7.

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compound of Claim 2.

18. A method of selectively increasing histamine levels in cells by contacting the cells with an antagonist of the histamine H3 receptor, said antagonists being a compound of Claim 9.

20 19 A method of selectively increasing histamine levels in cells by contacting the cells with an antagonist of the histamine H3 receptor, said antagonists being a compound of Claim 11.

 The method of Claim 15 wherein the antagonist is characterized by having little or no binding affinity for the histamine receptor H4R. A method for treatment or prevention of obesity which comprises administering to
a subject in need of such treatment or prevention an effective amount of a
compound of any of Claims 1-14.

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22. A method for treatment or prevention of a disorder or disease in which inhibition of the histamine H3 receptor has a beneficial effect which compnises administering to a subject in need of such treatment or prevention an effective amount of a compound of any of claims 1-14.

23. A method for treatment or prevention of a disorder or disease in which inhibition of the histamine H3 receptor has a beneficial effect which comprises administering to a subject in need of such treatment or prevention an effective amount of a compound of Claim 2.

24. A method for treatment or prevention of a disorder or disease in which inhibition of the histamine H3 receptor has a beneficial effect which comprises administering to a subject in need of such treatment or prevention an effective amount of a compound of Claim 7.

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25. A method for treatment or prevention of a disorder or disease in which inhibition of the histamine H3 receptor has a beneficial effect which comprises administering to a subject in need of such treatment or prevention an effective amount of a compound of Claim 9.

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26. A method for treatment or prevention of a disorder or disease in which inhibition of the histamine H3 receptor has a beneficial effect which comprises administering to a subject in need of such treatment or prevention an effective amount of a compound of Claim 11.

(12) INTERNATIONAL APPLICATION PUBLISHED UNDER THE PATENT COOPERATION TREATY (PCT)

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(3) International Publication Date 3 October 2002 (03.10.2002)

PCT

51) International Patent Chastideation¹: CO7C 217/58. Acid K 11792, Airlis, Acid P 200, 5200, COTO 25548, 254171, Acid P 200, 5200, COTO 25548, 254171, Acid P 2014, 21174, 21174, 21174, 21174, 21174, 21174, 21174, 21174, 21174, 21174, 21174, 21174, 21174, 21174, 21174, 21175, 21709, 21176, 21170, 211 217:00, 213:00) (51)

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Inventoria Applicants (for US out); BEAVERS, Lisa, Setson (USUS); 191 West State Road 22, Franklin, IN 46131 (US), GADSKI, Robert, ARB (USUS); 491 North Illinois, Indianapolis, IN 46238 (US), HIPSKININ.

alkylamine compounds of Formula (I) or pharmaceutically acceptable salts thereofwhich have selective compositions comprising such cyclic amines as well as methods (57) Abstract: The present invertion discloses novel substituted any histamine-H3 receptor antago activity as well as methods embodiment, invention discloses pharma

using them to treat

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Philip, Arrhur [USVUS]: 4255 South Cabin Court, New Philip, Arrhur [USVUS]: 4255 South Cabin Court, New Placistine, N. 4045 (US). LUNDSLEY, Craig, William [USVUS]: 252 East Lowell (US). LOBB, Kurne, Lyan [USVUS]: 562 East Lowell Arrhur, Indiampolit, N. 46219 (US). NIXON, James, Arthur [USVUS]: 737 Tao; Trail, Indiampolit, N. 46219 (US). STORM, Parthur [USVUS]: 129 Raintee Dive. Zionaville, N. 46077 (US). STAKARUWA, Takako [IPVUS]: 5919 Suscape Carle, Apdument BIST, Indiampolit, IN 46237 (US). WATSOW, Brian, Morgan [USVUS]: 3819 Suscape Carle, Apdument BIST, Indiampolit, IN 46237 (US). WATSOW, Brian, Morgan [USVUS]: 3816 Brian Place, Carmel, IN 46033 (US). (10) International Publication Number WO 02/076925 A3

Agents: WOOD, Dan, L. et al.; Bli Lilly And Company, P. O. Box 6288, Indianapolis, IN 46206-6288 (US). €

4) Designated States (regional): ARIPO patent (GH, GM, RE, LS, Mr. Uo, ZM, ZW), Bursain patent (AM, AZ, BY, KG, KZ, MD, RU, Ti, TM, Bunopean patent (AT, BE, CH, CY, DE, DK, ES, FI, FK, GB, GR, ET, TL, MC, NL, PT, SE, TR), OAPI patent (RF, BI, CT, CO, CT, CM, GA, GN, GO, GW, ML, MR, NE, TD, TG). 3

[Continued on next page]

(54) TIUR: NON-IMIDAZOLE ARYL ALKYLAMINES COMPOUNDS AS HISTAMINE H3 RECEPTOR ANTAGONISTS PREPARATION AND THERAPEUTIC USES

A3 WO 02/076925

Declarations under Rule 4.17:

MJ. MD. MG. MK. MD! MRT. MX. MZ. NO. NZ. OM, PH.
P. F. RO, NY. SD. SE, SG. S. SK. ST. TH, TN. RR.
TT. TZ. Ld., UG. UZ. NY. YU. ZJ. ZM, ZM, ZM, ARIPO patent
(GH. GM. KE. LS. MH, MZ. SD. SL, SZ. TZ. UG. ZM, ZW),
Eurazian patent (M. M. Z. B. KG. KZ. MD. RU. TJ. TJ.),
Eurapean patent (M. R. E. CH. CY. DE. DK. ES. FI, R. GB.
GR. ET. LU, MC. NL. PT. SE. TR), CMPI patent (BF. BJ.
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Published:

- with international search report

Date of publication of the international search report: 18 September 2003

For two-letter codes and other abbreviations, refer to the "Guid-ance Notes on Codes and Abbreviations" appearing at the begin-ning of each regular issue of the PCT Gazette.

INTERNATIONAL SEARCH REPORT

Int. Const Application No PCT/US 62/96644

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Minimum documentation securing (classification synthols)
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Further documents are listed in the continuation of box C.	* Special ontegories of ched documents ;	"A" document defining the general state of the art which is not considered to be of particular relevance

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Contraction of the Contract to	 A document defining the general state of the art which is not considered to be of particular retiveance 	E earlier document but published on or after the International filing date	L' do aument which may throw doubts on priority claim(s) or which is doed to establish the publication date of arruther orbital strategies on a resulted to the second of the properties of the	O' document referring to an oral disclosure, use, exhibition or other manner.	the man and the price of the balance that the price of th

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INTERNATIONAL SEARCH REPORT

mornational application No. PCT/US 02/06644

X	Observations where certain claims were found unsearchable (Continuation of item 1 of first sheet)	
E .	This international Search Report has not been established in respect of certain claims under Article 17(2)(a) for the following reasons:	
<u> </u>	Colons Nos. Decimie they natus to achied matter not required to be searched by this Authority, namely. Although claims 21-26 are directed to a method of treatment of the human/animal body, the search has been carried out and based on the alleged effects of the compounds.	
	Colors No.: because they relate to parts of the international Application that to not comply with the prescribed requirements to such an extent that no meaningful international Search can be carried out, specifically; see FURTHER INFORMATION sheet PCT/ISA/210	
	Claims Nos.; because they are dependent claims and are not drafted in accordance with the second and third sentences of Rule 6.4(s).	
Box II	Observations where unity of invention is lacking (Continuation of Item 2 of first sheet)].
T ster	This International Searching Authority found multiple inventions in this International application, as follows:	1
	see additional sheet	
<u>-</u>	As all required additional search foca were timely paid by the applicant, this international Search Report covers all searchable chalms.	
	As all searchable claims could be searched without effort justifying an additional lee, this Authority did not britte payment of any additional he.	
	As only some of the required additional esserb less were timely paid by the applicant, this international Gearch Raport covers only those claims for which less were paid, specifically deline. Nos.:	
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<u>-</u>	No required additional search fees were timely poid by the applicant. Corsequently, this international Search Report is restricted to the invention little metioned in the claims; it is covered by claims float.	
	1,2,4,7,14-17,20-24 all in part	
Remark	Remark to Drottest The additional search fees were accompanied by the applicant's protest.	
	No protest accompanied the payment of additional search fees.	
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ternational Application No. PCT/US 62 / 05644

FURTHER INFORMATION CONTINUED FROM PCT/ISA/ 210

This international Searching Authority found multiple (groups of) inventions in this international application, as follows:

1. Claims: 1,2,4,7,14-17,20-24 all in part

Benzene compounds of general formulas 1 or 11 with R6 = hydrogen or halo and X = 0xygen, compositions and methods using these compounds.

2. Claims: 1-4,6,7,14-17,20-24 in part, 8,9,11,18,19,25,26

Benzene compounds of general formulas I or II with R6 = hydrogen or halo and X = N or NR7, compositions and methods using these compounds.

3. Claims: 1,2,4,7,14-17,20-24 all in part

Benzene compounds of general formulas I or II with R6 = hydrogen or halo and \hat{X} = sulfur, compositions and methods using these compounds.

4. Claims: 1-3,6,7,14-17,20-24 all in part

Carbobicyclic compounds of general formulas I or II with R6 cyclized with the attached carbon atom at the R5 position, compositions and methods using these compounds.

5. Claims: 1-3,6,7,14-17,20-24 in part, 5,10,12,13

Tetrahydroisoquinoline compounds of general formulas I or II with R6 cyclized with the attached carbon atom at the R7 position; compositions and methods using these compounds.

International Application No. PCT/US 92/96644

FURTHER INFORMATION CONTINUED FROM PCT/ISA/ 219

Continuation of Box 1.2

The initial phase of the search for invention I revealed a very large number of documents relevant to the issue of novelty. So many documents were retrieved that it is impossible to determine which parts of the claims may be said to define subject-matter for which protection might legitimately be sought (Article 5 PCT). For these reasons, a meaningful the search over the whole breadth of the claims is impossible. Consequently, the search for invention I has been restricted to the compounds of the examples.

The applicant's attention is drawn to the fact that claims, or parts of claims, relating to inventions in respect of which no international. Search report has been established need not be the subject of an international preliminary examination (Rule 66.1(e) PCT). The applicant is advised that the EPO policy when acting as an International Preliminary Examining Authority is normally not to carry out a preliminary examination on matter which has not been searched. This is the case irrespective of whether or not the claims are amended following receipt of the search report or during any Chapter II procedure.

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